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6. ALPHA., 9. ALPHA.- DIFLUORO-17. ALPHA.- (2- FURANYLCARBOXYL) OXY- 11. BETA.- HYDROXY-16. ALPHA.- METHYL-3-OXO-ANDROST-1,4-DIENE-17-CARBOTHIOIC ACID S-FLUOROMETHYL ESTER ALS ENTZÜNDUNGSHEMMENDES ARZNEIMITTEL

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The present invention relates to a novel anti-inflammatory and anti-allergic compound of the androstane series and to processes for its preparation. The present invention also relates to pharmaceutical formulations containing the compound and to therapeutic uses thereof, particularly for the treatment of inflammatory and allergic conditions.

Glucocorticoids which have anti-inflammatory properties are known and are widely used for the treatment of inflammatory disorders or diseases such as asthma and rhinitis. For example, US Patent 4335121 discloses 6α, 9α-Difluoro-17α-(1-oxopropoxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (known by the generic name of fluticasone propionate) and derivatives thereof. The use of glucocorticoids generally, and especially in children, has been limited in some quarters by concerns over potential side effects. The side effects that are feared with glucocorticoids include suppression of the Hypothalamic-Pituitary-Adrenal (HPA) axis, effects on bone growth in children and on bone density in the elderly, ocular complications (cataract formation and glaucoma) and skin atrophy. Certain glucocorticoid compounds also have complex paths of metabolism wherein the production of active metabolites may make the pharmacodynamics and pharmacokinetics of such compounds difficult to understand. Whilst the modern steroids are very much safer than those originally introduced, it remains an object of research to produce new molecules which have excellent anti-inflammatory properties, with predictable pharmacokinetic and pharmacodynamic properties, with an attractive side effect profile, and with a convenient treatment regime.

We have now identified a novel glucocorticoid compound which substantially meets these objectives.

Thus, according to one aspect of the invention, there is provided a compound of formula (I)

and solvates thereof.

The chemical name of the compound of formula (I) is 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester.

References hereinafter to the compound according to the invention include both the compound of formula (I) and solvates thereof, particularly pharmaceutically acceptable solvates.

The compound of formula (I) has potentially beneficial anti-inflammatory or anti-allergic effects, particularly upon topical administration, demonstrated by, for example, its ability to bind to the glucocorticoid receptor and to illicit a response via that receptor. Hence, the compound of formula (I) is useful in the treatment of inflammatory and/or allergic disorders.

Compound (I) undergoes highly efficient hepatic metabolism to yield the 17-β carboxylic acid (X) as the sole major metabolite in rat and human in vitro systems. This metabolite has been synthesised and demonstrated to be >1000 fold less active than the parent compound in in vitro functional glucocorticoid assays.
This efficient hepatic metabolism is reflected by in vivo data in the rat, which have demonstrated plasma clearance at a rate approaching hepatic blood flow and an oral bioavailability of <1%, consistent with extensive first-pass metabolism.

In vitro metabolism studies in human hepatocytes have demonstrated that compound (I) is metabolised in an identical manner to fluticasone propionate but that conversion of (I) to the inactive acid metabolite occurs approximately 5-fold more rapidly than with fluticasone propionate. This very efficient hepatic inactivation would be expected to minimise systemic exposure in man leading to an improved safety profile.

Inhaled steroids are also absorbed through the lung and this route of absorption makes a significant contribution to systemic exposure. Reduced lung absorption could therefore provide an improved safety profile. Studies with compound of formula (I) have shown significantly lower exposure to compound of formula (I) than with fluticasone propionate after dry powder delivery to the lungs of anaesthetised pigs.

An improved safety profile is believed to allow the compound of formula (I) to demonstrate the desired anti-inflammatory effects when administered once-per day. Once-per-day dosing is considered to be significantly more convenient to patients than the twice-per day dosing regime that is normally employed for fluticasone propionate.

Examples of disease states in which the compound of the invention has utility include skin diseases such as eczema, psoriasis, allergic dermatitis, neurodermatitis, pruritis and hypersensitivity reactions; inflammatory conditions of the nose, throat or lungs such as asthma (including allergen-induced asthmatic reactions), rhinitis (including hayfever), nasal polyps, chronic obstructive pulmonary disease, interstitial lung disease, and fibrosis; inflammatory bowel conditions such as ulcerative colitis and Crohn’s disease; and auto-immune diseases such as rheumatoid arthritis.

The compound of the invention may also have use in the treatment of conjunctiva and conjunctivitis.

It will be appreciated by those skilled in the art that reference herein to treatment extends to prophylaxis as well as the treatment of established conditions.

As mentioned above, the compound of formula (I) is useful in human or veterinary medicine, in particular as an anti-inflammatory and anti-allergic agent.

There is thus provided as a further aspect of the invention the compound of formula (I) or a physiologically acceptable solvate thereof for use in human or veterinary medicine, particularly in the treatment of patients with inflammatory and/or allergic conditions, especially for treatment once-per-day.

According to another aspect of the invention, there is provided the use of the compound of formula (I) or physiologically acceptable solvate thereof for the manufacture of a medicament for the treatment of patients with inflammatory and/or allergic conditions, especially for treatment once-per-day.

In a further or alternative aspect, there is provided a method for the treatment of a human or animal subject with an inflammatory and/or allergic condition, which method comprises administering to said human or animal subject an effective amount of the compound of formula (I) or physiologically acceptable solvate thereof, especially for administration once-per-day.

The compound according to the invention may be formulated for administration in any convenient way, and the invention therefore also includes within its scope pharmaceutical compositions comprising the compound of formula (I) or a physiologically acceptable solvate thereof together, if desirable, in admixture with one or more physiologically acceptable diluents or carriers. Pharmaceutical compositions suitable for once-per-day administration are of particular interest.

Further, there is provided a process for the preparation of such pharmaceutical compositions which comprises mixing the ingredients.

The compound according to the invention may, for example, be formulated for oral, buccal, sublingual, parenteral, local or rectal administration, especially local administration.

Local administration as used herein, includes administration by insufflation and inhalation. Examples of various types of preparation for local administration include ointments, lotions, creams, gels, foams, preparations for delivery by transdermal patches, powders, sprays, aerosols, capsules or cartridges for use in an inhaler or insufflator or drops (eg eye or nose drops), solutions/suspensions for nebulisation, suppositories, pessaries, retention enemas and chewable or suckable tablets or pellets (eg for the treatment of aphthous ulcers) or liposome or microencapsulation preparations.

Advantageously compositions for topical administration to the lung include dry powder compositions and spray compositions.

Dry powder compositions for topical delivery to the lung may, for example, be presented in capsules and cartridges for use in an inhaler or insufflator of, for example, gelatine. Formulations generally contain a powder mix for inhalation of the compound of the invention and a suitable powder base such as lactose or starch. Each capsule or cartridge may generally contain between 20μg-10mg of the compound of formula (I). Alternatively, the compound of the invention may be presented without excipients. Packaging of the formulation may be suitable for unit dose or multi-dose delivery. In the case of multi-dose delivery, the formulation can be pre-metered (eg as in Diskus, see GB 2242134 or Diskhaler, see GB 2178965, 2129691 and 2169265) or metered in use (eg as in Turbuhaler, see EP 69715). An example of a unit-dose device is Rotahaler (see GB 2064336). The Diskus inhalation device comprises an elongate strip formed
from a base sheet having a plurality of recesses spaced along its length and a lid sheet hermetically but peelably sealed thereto to define a plurality of containers, each container having therein an inhalable formulation containing a compound of formula (I) preferably combined with lactose. Preferably, the strip is sufficiently flexible to be wound into a roll. The lid sheet and base sheet will preferably have leading end portions which are not sealed to one another and at least one of the said leading end portions is constructed to be attached to a winding means. Also, preferably the hermetic seal between the base and lid sheets extends over their whole width. The lid sheet may preferably be peeled from the base sheet in a longitudinal direction from a first end of the said base sheet.

[0026] Pharmaceutical formulations which are non-pressurised and adapted to be administered as a dry powder sheet in a longitudinal direction from a first end of the said base sheet.

[0027] Spray compositions may for example be formulated as aqueous solutions or suspensions or as aerosols delivered from pressurised packs, such as a metered dose inhaler, with the use of a suitable liquefied propellant. Aerosol compositions suitable for inhalation such as can be either a suspension or a solution and generally contain the compound of formula (I) and a suitable propellant such as a fluorocarbon or hydrogen-containing chlorofluorocarbon or mixtures thereof, particularly hydrofluoroalkanes, especially 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoro-n-propene or a mixture thereof. The aerosol composition may optionally contain additional formulation excipients well known in the art such as surfactants eg oleic acid or lecithin and cosolvents eg ethanol. One example formulation is excipient free and consists essentially of (eg consists of) compound of formula (I) (preferably in unsolvated form eg as Form 1) (optionally in combination with another therapeutically active ingredient) and a propellant selected from 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoro-n-propene and mixture thereof. Another example formulation comprises particulate compound of formula (I), a propellant selected from 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoro-n-propene and mixture thereof and a suspending agent which is soluble in the propellant eg an oligolactic acid or derivative thereof as described in WO94/21229. The preferred propellant is 1,1,1,2-tetrafluoroethane. As noted elsewhere in this specification, compound of formula (I) does not appear to form a solvate with 1,1,1,2-tetrafluoroethane. Pressurised formulations will generally be retained in a canister (eg an aluminium canister) closed with a valve (eg a metering valve) and fitted into an actuator provided with a mouthpiece.

[0028] Pressurised aerosol formulations preferably do not comprise particulate medicament, a propellant and a stabiliser comprising a water addition (i.e. water added in addition to nascent formulation water). Pressurised aerosol formulations also preferably do not comprise particulate medicament, a propellant and a stabiliser comprising an amino acid, a derivative thereof or a mixture thereof.

[0029] Medicaments for administration by inhalation desirably have a controlled particle size. The optimum particle size for inhalation into the bronchial system is usually 1-10 μm, preferably 2-5 μm. Particles having a size above 20 μm are generally too large when inhaled to reach the small airways. To achieve these particle sizes the particles of compound of formula (I) as produced may be size reduced by conventional means eg by micronisation. The desired fraction may be separated out by air classification or sieving. Preferably, the particles will be crystalline, prepared for example by a process which comprises mixing in a continuous flow cell in the presence of ultrasonic radiation a flowing solution of compound of formula (I) as medicament in a liquid solvent with a flowing liquid antisolvent for said medicament (eg as described in International Patent Application PCT/GB99/04368) or else by a process which comprises admitting a stream of solution of the substance in a liquid solvent and a stream of liquid antisolvent for said substance tangentially into a cylindrical mixing chamber having an axial outlet port such that said streams are thereby intimately mixed through formation of a vortex and precipitation of crystalline particles of the substance is thereby caused (eg as described in International Patent Application PCT/GB00/04237). When an excipient such as lactose is employed, generally, the particle size of the excipient will be much greater than the inhaled medicament within the present invention. When the excipient is lactose it will typically be present as milled lactose, wherein not more than 85% of lactose particles will have a MMD of 60-90 μm and not less than 15% will have a MMD of less than 15 μm.

[0030] Formulations for administration topically to the nose (eg for the treatment of rhinitis) include pressurised aerosol formulations and aqueous formulations administered to the nose by pressurised pump. Formulations which are non-pressurised and adapted to be administered topically to the nasal cavity are of particular interest. The formulation preferably contains water as the diluent or carrier for this purpose. Aqueous formulations for administration to the lung or nose may be provided with conventional excipients such as buffering agents, tonicity modifying agents and the like. Aqueous formulations may also be administered to the nose by nebulisation.

[0031] Other possible presentations include the following:

[0032] Ointments, creams and gels, may, for example, be formulated with an aqueous or oily base with the addition of suitable thickening and/or gelling agent and/or solvents. Such bases may thus, for example, include water and/or an oil such as liquid paraffin or a vegetable oil such as arachis oil or castor oil, or a solvent such as polyethylene glycol. Thickening agents and gelling agents which may be used according to the nature of the base include soft paraffin, aluminium stearate, cetostearyl alcohol, polyethylene glycols, woolfat, beeswax, carboxypolymethylene and cellulose derivatives, and/or glyceryl monostearate and/or non-ionic emulsifying agents.
Lotions may be formulated with an aqueous or oily base and will in general also contain one or more emulsifying agents, stabilising agents, dispersing agents, suspending agents or thickening agents.

Powders for external application may be formed with the aid of any suitable powder base, for example, talc, lactose or starch. Drops may be formulated with an aqueous or non-aqueous base also comprising one or more dispersing agents, solubilising agents, suspending agents or preservatives.

If appropriate, the formulations of the invention may be buffered by the addition of suitable buffering agents.

The proportion of the active compound of formula (I) in the local compositions according to the invention depends on the precise type of formulation to be prepared but will generally be within the range of from 0.001 to 10% by weight. Generally, however for most types of preparations advantageously the proportion used will be within the range of from 0.005 to 1% and preferably 0.01 to 0.5%. However, in powders for inhalation or insufflation the proportion used will usually be within the range of from 0.1 to 5%.

Aerosol formulations are preferably arranged so that each metered dose or "puff" of aerosol contains 1µg-2000µg eg 20µg-2000µg, preferably about 20µg-500µg of a compound of formula (I). Administration may be once daily or several times daily, for example 2, 3, 4 or 8 times, giving for example 1, 2 or 3 doses each time. Preferably the compound of formula (I) is delivered once or twice daily, more preferably once per day. The overall daily dose with an aerosol will typically be within the range 10µg-10mg eg 100µg-10mg preferably, 200µg-2000µg.

Topical preparations may be administered by one or more applications per day to the affected area; over skin areas occlusive dressings may advantageously be used. Continuous or prolonged delivery may be achieved by an adhesive reservoir system.

For internal administration the compound according to the invention may, for example, be formulated in conventional manner for oral, parenteral or rectal administration. Formulations for oral administration include syrups, elixirs, powders, granules, tablets and capsules which typically contain conventional excipients such as binding agents, fillers, lubricants, disintegrants, wetting agents, suspending agents, emulsifying agents, preservatives, buffer salts, flavouring, colouring and/or sweetening agents as appropriate. Dosage unit forms are, however, preferred as described below.

Preferred forms of preparation for internal administration are dosage unit forms i.e. tablets and capsules. Such dosage unit forms contain from 0.1mg to 20mg preferably from 2.5 to 10mg of the compound of the invention.

The compound according to the invention may in general be given by internal administration in cases where systemic adreno-cortical therapy is indicated.

In general terms preparations, for internal administration may contain from 0.05 to 10% of the active ingredient dependent upon the type of preparation involved. The daily dose may vary from 0.1mg to 60mg, eg 5-30mg, dependent on the condition being treated, and the duration of treatment desired.

Slow release or enteric coated formulations may be advantageous, particularly for the treatment of inflammatory bowel disorders.

The pharmaceutical compositions according to the invention may also be used in combination with another therapeutically active agent, for example, a β2-adrenoreceptor agonist, an anti-histamine or an anti-allergic. The invention thus provides, in a further aspect, a combination comprising the compound of formula (I) or a physiologically acceptable solvate thereof together with another therapeutically active agent, for example, a β2-adrenoreceptor agonist, an anti-histamine or an anti-allergic.

Examples of β2-adrenoreceptor agonists include salmeterol (eg as racemate or a single enantiomer such as the R-enantiomer), salbutamol, formoterol, salmefamol, fenoterol or terbutaline and salts thereof, for example the xinafoate salt of salmeterol, the sulphate salt or free base of salbutamol or the fumarate salt of formoterol. Examples of anti-histamines include methapyrilene or loratadine.

Other suitable combinations include, for example, other anti-inflammatory agents eg NSAIDs (eg sodium cromoglicate, nedocromil sodium, PDE4 inhibitors, leukotriene antagonists, iNOS inhibitors, tryptase and elastase inhibitors, beta-2 integrin antagonists and adenosine 2a agonists)) or antiinfective agents (eg antibiotics, antivirals).

Of particular interest is use of the compound of formula (I) in combination with a phosphodiesterase 4 (PDE4) inhibitor. The PDE4-specific inhibitor useful in this aspect of the invention may be any compound that is known to inhibit the PDE4 enzyme or which is discovered to act as a PDE4 inhibitor, and which are only PDE4 inhibitors, not compounds which inhibit other members of the PDE family as well as PDE4. Generally it is preferred to use a PDE4 inhibitor which has an IC50 ratio of about 0.1 or greater as regards the IC50 for the PDE4 catalytic form which binds rolipram with a high affinity divided by the IC50 for the form which binds rolipram with a low affinity. For the purposes of this disclosure, the cAMP catalytic site which binds R and S rolipram with a low affinity is denominated the "low affinity" binding site (LPDE 4) and the other form of this catalytic site which binds rolipram with a high affinity is denominated the "high affinity" binding site (HPDE 4). This term "HPDE4" should not be confused with the term "hPDE4" which is used to denote human PDE4. Initial experiments were conducted to establish and validate a [3H]-rolipram binding assay. Details of this work are given in the Binding Assays described in detail below.

The preferred PDE4 inhibitors of use in this invention will be those compounds which have a salutary therapeutic ratio, i.e., compounds which preferentially inhibit cAMP catalytic activity where the enzyme is in the form that binds...
rolipram with a low affinity, thereby reducing the side effects which apparently are linked to inhibiting the form which binds rolipram with a high affinity. Another way to state this is that the preferred compounds will have an IC$_{50}$ ratio of about 0.1 or greater as regards the IC$_{50}$ for the PDE4 catalytic form which binds rolipram with a high affinity divided by the IC$_{50}$ for the form which binds rolipram with a low affinity.

A further refinement of this standard is that of one wherein the PDE4 inhibitor has an IC$_{50}$ ratio of about 0.1 or greater; said ratio is the ratio of the IC$_{50}$ value for competing with the binding of 1nM of [3H]R-rolipram to a form of PDE4 which binds rolipram with a high affinity divided by the IC$_{50}$ value for inhibiting the PDE4 catalytic activity of a form which binds rolipram with a low affinity using 1 μM[3H]-cAMP as the substrate.

**[0049]** Examples of useful PDE4 inhibitors are:

- (R)-(+)-1-(4-bromobenzyl)-4-[(3-cyclopentyloxy)-4-methoxyphenyl]-2-pyrrolidone;
- (R)-(+)-1-(4-bromobenzyl)-4-[(3-cyclopentyloxy)-4-methoxyphenyl]-2-pyrrolidone;
- 3-(cyclopentyloxy-4-methoxyphenyl)-1-(4-N'-[N2-cyano-S-methyl-isothioureido]benzyl)-2-pyrrolidone;
- cis 4-cyano-4-[(3-cyclopentyloxy-4-methoxyphenyl)cyclohexan-1-carboxylic acid];
- cis-[4-cyano-4-(3-cyclopropylmethoxy-4-difluoromethoxyphenyl)cyclohexan-1-ol];
- (R)-(+-)ethyl [4-[(3-cyclopentyloxy-4-methoxyphenyl)pyrrolidine-2-ylidene]acetate; and
- (S)-(--)ethyl [4-[(3-cyclopentyloxy-4-methoxyphenyl)pyrrolidine-2-ylidene]acetate.

**[0050]** Most preferred are those PDE4 inhibitors which have an IC$_{50}$ ratio of greater than 0.5, and particularly those compounds having a ratio of greater than 1.0. Preferred compounds are cis 4-cyano-4-(3-cyclopentyloxy-4-methoxyphenyl)cyclohexan-1-carboxylic acid, 2-carbomethoxy-4-cyano-4-(3-cyclopropylmethoxy-4-difluoromethoxyphenyl)cyclohexan-1-one and cis-[4-cyano-4-(3-cyclopropylmethoxy-4-difluoromethoxyphenyl)cyclohexan-1-ol]; these are examples of compounds which bind preferentially to the low affinity binding site and which have an IC$_{50}$ ratio of 0.1 or greater. Other compounds of interest include:

- Compounds set out in U.S. patent 5,552,438 issued 03 September, 1996; this patent and the compounds it discloses are incorporated herein in full by reference. The compound of particular interest, which is disclosed in U.S. patent 5,552,438, is cis-[4-cyano-4-(3-cyclopentyloxy-4-methoxyphenyl)cyclohexan-1-carboxylic acid (also known as cimolast) and its salts, esters, pro-drugs or physical forms;

### Phosphodiesterase and Rolipram Binding Assays

**Assay method 1A**

- Isolated human monocyte PDE4 and hrPDE (human recombinant PDE4) was determined to exist primarily in the low affinity form. Hence, the activity of test compounds against the low affinity form of PDE4 can be assessed using standard assays for PDE4 catalytic activity employing 1 μM [3H]cAMP as a substrate (Torphy et al., J. of Biol. Chem., Vol. 267, No. 3 pp1798-1804, 1992).

**Assay method 1B**

- Measurement of Phosphodiesterase Activity

- PDE activity was assayed using a [3H]cAMP SPA or [3H]cGMP SPA enzyme assay as described by the supplier.
(Amersham Life Sciences). The reactions were conducted in 96-well plates at room temperature, in 0.1 ml of reaction buffer containing (final concentrations): 50 mM Tris-HCl, pH 7.5, 8.3 mM MgCl₂, 1.7 mM EGTA, [³H]cAMP or [³H] cGMP (approximately 2000 dpm/pmol), enzyme and various concentrations of the inhibitors. The assay was allowed to proceed for 1 hr and was terminated by adding 50 μl of SPA yttrium silicate beads in the presence of zinc sulfate. The plates were shaken and allowed to stand at room temperature for 20 min. Radiolabeled product formation was assessed by scintillation spectrometry.

[³H]R-rolipram binding assay

[0053] The [³H]R-rolipram binding assay was performed by modification of the method of Schneider and co-workers, see Nicholson, et al., Trends Pharmacol. Sci., Vol. 12, pp.19-27 (1991) and McHale et al., Mol. Pharmacol., Vol. 39, 109-113 (1991). R-Rolipram binds to the catalytic site of PDE4 see Torphy et al., Mol. Pharmacol., Vol. 39, pp. 376-384 (1991). Consequently, competition for [³H]R-rolipram binding provides an independent confirmation of the PDE4 inhibitor potencies of unlabeled competitors. The assay was performed at 30°C for 1 hr in 0.5 μl buffer containing (final concentrations): 50 mM Tris-HCl, pH 7.5, 5 mM MgCl₂, 0.05% bovine serum albumin, 2 nM [³H]R-rolipram (5.7 x 10⁴ dpm/pmol) and various concentrations of non-radiolabeled inhibitors. The reaction was stopped by the addition of 2.5 ml of ice-cold reaction buffer (without [³H]R-rolipram) and rapid vacuum filtration (Brandel Cell Harvester) through Whatman GF/B filters that had been soaked in 0.3% polyethylenimine. The filters were washed with an additional 7.5 ml of cold buffer, dried, and counted via liquid scintillation spectrometry.

[0054] The invention thus provides, in a further aspect, a combination comprising the compound of formula (I) or a physiologically acceptable solvate thereof together with a PDE4 inhibitor.

[0055] The combination referred to above may conveniently be presented for use in the form of a pharmaceutical formulation and thus pharmaceutical formulations comprising a combination as defined above together with a physiologically acceptable diluent or carrier represent a further aspect of the invention.

[0056] The individual compounds of such combinations may be administered either sequentially or simultaneously in separate or combined pharmaceutical formulations.

[0057] Appropriate doses of known therapeutic agents will be readily appreciated by those skilled in the art.

[0058] Surprisingly, the compound of formula (I) has demonstrated a significant propensity to form solvates with commonly used organic solvents. Such solvates are essentially stoichiometric eg the ratio of compound of formula (I) to solvent is close to 1:1 eg according to Applicant’s analysis has been determined to be in the range 0.95-1.05: 1. For example, we have prepared solvates with solvents such as acetone, dimethylformamide (DMF), dimethylacetamide (DMAc), tetrahydrofuran (THF), N-methyl-2-pyrrolidone, isopropanol and methylisobutylketone. The solvation of compound of formula (I) is not predictable however since we have found that even though it does form a solvate with isopropanol it does not appear to form a solvate with ethanol or methanol. Furthermore it does not appear to form a solvate with 1,1,1,2-tetrafluoroethane, ethylacetate, methylacetate, toluene, methylisobutylketone (MIBK) or water either. However due to the toxicity of many organic solvents it has been necessary to develop special final stage processing conditions (discussed later) in order to permit the compound of formula (I) to be produced in unsolvated form. Thus according to another aspect of the invention there is provided a compound of formula (I) in unsolvated form.

[0059] Surprisingly we have also discovered that the compound of formula (I) in unsolvated form may exist in a number of polymorphic forms. Specifically we have identified polymorphic forms which may be distinguished by means of X-Ray Powder Diffraction (XRPD) which we have named as Form 1, Form 2 and Form 3. Form 3 appears to be an unstable minor polymorphic modification of Form 2. Broadly speaking the Forms are characterised in their XRPD profiles as follows:

Form 1: Peak at around 18.9 degrees 2Theta
Form 2: Peaks at around 18.4 and 21.5 degrees 2Theta.
Form 3: Peaks at around 18.6 and 19.2 degrees 2Theta.

[0060] Within the range 21-23 degrees 2Theta Form 3 shows a single peak whereas Form 2 shows a pair of peaks. A peak at 7 degrees 2Theta is present in all cases however it is present at much higher intensity in the case of Forms 2 and 3 than is the case for Form 1.

[0061] The XRPD patterns of the polymorphs are shown overlaid in Figure 1. The conversion of Form 2 to Form 1 with time in an aqueous slurry at ambient temperature is shown in Figure 2. In the conversion of Form 2 to Form 1 the loss of a peak characteristic of Form 2 (labelled B) at around 18.4 degrees 2Theta, a marked reduction in intensity in the peak at around 7 degrees 2Theta (labelled A) and the appearance of a peak characteristic of Form 1 (labelled C) at around 18.9 degrees 2Theta are particularly noticeable.

[0062] The temperature dependence of Form 3 is shown in Figure 4. The temperature was varied according to the profile shown in Figure 5. From Figure 4 it can be seen that Form 3 converts first to Form 2 over the temperature range 30-170 °C and then converts to Form 1 over the temperature range 170-230 °C. In the conversion of Form 3 to Form 2
the division of one peak in the range 21-23 degrees 2Theta into two peaks within the same range and the shifting leftwards of the peak at around 18.6 degrees 2Theta to around 18.4 degrees 2Theta are particularly noticeable. In the conversion of Form 2 to Form 1 similar changes to those noted in the previous paragraph may be observed.

[0063] The differential scanning calorimetry (DSC) and thermal gravimetric analysis (TGA) profiles of Form 1 are shown in Figure 3. The profiles are characterised by a transition at around 280-300 °C (typically close to 298 °C) corresponding to an endothermic event in the DSC and chemical degradation in the TGA. The DSC profiles of Forms 2 and 3 were not materially different under the conditions of the experiments performed and thus DSC is not a suitable technique for distinguishing between the 3 Forms. In Figure 3 the absence of activity in the TGA and DSC profiles below around 298 °C implies that the substance shows good physical and chemical stability at normal operating temperatures.

[0064] As shown in the Examples, enthalpy of dissolution of Forms 1 and 3 have been determined in certain organic solvents and accordingly an enthalpy of transition from Form 3 to Form 1 of 5.1-6.7 kJ/mol has been estimated.

[0065] Thus we prefer compound of formula (I) in unsolvated Form 1 since this form appears to be thermodynamically most stable at ambient temperature and also appears to be least susceptible to undesirable moisture sorption (see results in Examples section). Nevertheless Form 2 (or Form 3) may be preferred under other conditions.

[0066] Although use of a compound of formula (I) in solvated form is not preferred, nevertheless we have surprisingly found that certain solvate forms have particularly attractive physicochemical properties which makes them useful as intermediates in the preparation of a compound of formula (I) in unsolvated form (eg by removal of solvent as a final step). For example we have discovered that certain stoichiometric solvates can be isolated as solids in highly crystalline form. Thus we also provide as an aspect of the invention:

- Compound of formula (I) as the methylethylketone solvate
- Compound of formula (I) as the isopropanol solvate
- Compound of formula (I) as the tetrahydrofuran solvate
- Compound of formula (I) as the acetone solvate.

In particular we provide the aforementioned solvates as solids in crystalline form. A further particular advantage of these solvates is the fact that desolvation of the solvate (eg by heating) results in formation of the unsolvated form as the preferred Form 1. The aforementioned solvates have relatively low toxicity and are suitable for use in industrial scale manufacture. Compound of formula (I) as the DMF solvate which may also be isolated as a solid in crystalline form is also of interest for use in onward processing to unsolvated Form 1.

[0067] The compound of formula (I) and solvates thereof may be prepared by the methodology described hereinafter, constituting a further aspect of this invention.

[0068] A process according to the invention for preparing a compound of formula (I) or a solvate thereof comprises alkylation of a thioacid of formula (II)

\[
\text{O} \quad \text{SH} \\
\text{O} \quad \text{C} \\
\text{HO} \quad \text{CH}_3 \\
\text{CH}_3 \\
\text{O} \\
\text{CH}_3 \\
\text{F} \quad \text{H} \quad \text{L} \\
\text{(II)}
\]

or a salt thereof.

[0069] In this process the compound of formula (II) may be reacted with a compound of formula FCH\_2L wherein L represents a leaving group (eg a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane.

[0070] As noted later, preferably the compound of formula (II) is employed as a salt, particularly the salt with diisopropylethylamine.

[0071] In a preferred process for preparing the compound of formula (I), the compound of formula (II) or a salt thereof is treated with bromofluoromethane optionally in the presence of a phase transfer catalyst. A preferred solvent is methylacetate, or more preferably ethylacetate, optionally in the presence of water. The presence of water improves solubility of both starting material and product and the use of a phase transfer catalyst results in an increased rate of reaction.
Examples of phase transfer catalysts that may be employed include (but are not restricted to) tetrabutylammonium bromide, tetrabutylammonium chloride, benzyltributylammonium bromide, benzyltributylammonium chloride, benzyltributyrammonium bromide, benzyltributyrammonium chloride, methyltributylammonium bromide and methyltributyrammonium chloride. THF has also successfully been employed as solvent for the reaction wherein the presence of a phase transfer catalyst again provides a significantly faster reaction rate. Preferably the product present in an organic phase is washed firstly with aqueous acid eg dilute HCl in order to remove amine compounds such as triethylamine and diisopropylethylamine and then with aqueous base eg sodium bicarbonate in order to remove any unreacted precursor compound of formula (II). As noted later, if the compound of formula (I) so produced in solution in ethylacetate is distilled and toluene added, then unsolvated Form 1 crystallises out.

[0072] Compounds of formula (II) may be prepared from the corresponding 17α-hydroxyl derivative of formula (III):

![Chemical Structure of III](image)

using for example, the methodology described by G. H. Phillipps et al., (1994) Journal of Medicinal Chemistry, 37, 3717-3729. For example the step typically comprises the addition of a reagent suitable for performing the esterification eg an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride (employed in at least 2 times molar quantity relative to the compound of formula (III)) in the presence of an organic base eg triethylamine. The second mole of 2-furoyl chloride reacts with the thioacid moiety in the compound of formula (III) and needs to be removed eg by reaction with an amine such as diethylamine.

[0073] This method suffers disadvantages, however, in that the resultant compound of formula (II) is not readily purified of contamination with the by-product 2-furoyldiethylamide. We have therefore invented several improved processes for performing this conversion.

[0074] In a first such improved process we have discovered that by using a more polar amine such as diethanolamine, a more water soluble by-product is obtained (in this case 2-furoyldiethanolamide) which permits compound of formula (II) or a salt thereof to be produced in high purity since the by-product can efficiently be removed by water washing.

[0075] Thus according to this aspect of the invention we provide a process for preparing a compound of formula (II) which comprises:

(a) reacting a compound of formula (III) with an activated derivative of 2-furoic acid as in an amount of at least 2 moles of the activated derivative per mole of compound of formula (III) to yield a compound of formula (IIA)

![Chemical Structure of IIA](image)

and

(b) removal of the sulphur-linked 2-furoyl moiety from compound of formula (IIA) by reaction of the product of step
In two particularly convenient embodiments of this process we also provide methods for the efficient purification of the end product which comprise either

(c1) when the product of step (b) is dissolved in a substantially water immiscible organic solvent, purifying the compound of formula (II) by washing out the amide by-product from step (b) with an aqueous wash, or
(c2) when the product of step (b) is dissolved in a water miscible solvent, purifying the compound of formula (II) by treating the product of step (b) with an aqueous medium so as to precipitate out pure compound of formula (II) or a salt thereof.

In step (a) preferably the activated derivative of 2-furoic acid may be an activated ester of 2-furoic acid, but is more preferably a 2-furoyl halide, especially 2-furoyl chloride. A suitable solvent for this reaction is ethylacetate or methylacetate (preferably methylacetate) (when step (c1) may be followed) or acetone (when step (c2) may be followed). Normally an organic base eg triethylamine will be present. In step (b) preferably the organic base is diethanolamine. The base may suitably be dissolved in a solvent eg methanol. Generally steps (a) and (b) will be performed at reduced temperature eg between 0 and 5 °C. In step (c1) the aqueous wash may be water, however the use of brine results in higher yields and is therefore preferred. In step (c) the aqueous medium is for example a dilute aqueous acid such as dilute HCl.

According to a related aspect of the invention we provide an alternative process for preparing a compound of formula (II) which comprises:

(a) reacting a compound of formula (III) with an activated derivative of 2-furoic acid in an amount of at least 2 moles of activated derivative per mole of compound of formula (III) to yield a compound of formula (IIA) ; and
(b) removal of the sulphur-linked 2-furoyl moiety from compound of formula (IIA) by reaction of the product of step (a) with a further mole of compound of formula (III) to give two moles of compound of formula (II).

In step (a) preferably the activated derivative of 2-furoic acid may be an activated ester of 2-furoic acid, but is more preferably a 2-furoyl halide, especially 2-furoyl chloride. A suitable solvent for this step is acetone. Normally an organic base eg triethylamine will be present. In step (b) a suitable solvent is DMF or dimethylacetamide. Normally an organic base eg triethylamine will be present. Generally steps (a) and (b) will be performed at reduced temperature eg between 0 and 5 °C. The product may be isolated by treatment with acid and washing with water.

This aforementioned process is very efficient in that it does not produce any furoylamide by-product (thus affording inter alia environmental advantages) since the excess mole of furoyl moiety is taken up by reaction with a further mole of compound of formula (II) to form an additional mole of compound of formula (II).

Further general conditions for the conversion of compound of formula (III) to compound of formula (II) in the two processes just described will be well known to persons skilled in the art.

According to a preferred set of conditions, however, we have found that the compound of formula (II) may advantageously be isolated in the form of a solid crystalline salt. The preferred salt is a salt formed with a base such as triethylamine, 2,4,6-trimethylpyridine, diisopropylethylamine or N-ethylpiperidine. Such salt forms of compound of formula (II) are more stable, more readily filtered and dried and can be isolated in higher purity than the free thioacid. The most preferred salt is the salt formed with diisopropylethylamine. The triethylamine salt is also of interest.

Compounds of formula (III) may be prepared in accordance with procedures described in GB 2088877B.

Compounds of formula (III) may also be prepared by a process comprising the following steps:

(a) with an organic primary or secondary amine base capable of forming a water soluble 2-furoyl amide.

Step (a) comprises oxidation of a solution containing the compound of formula (V). Preferably, step (a) will be performed in the presence of a solvent comprising methanol, water, tetrahydrofuran, dioxan or diethylene glycol dimethyl ether. So as to enhance yield and throughput, preferred solvents are methanol, water or tetrahydrofuran, and more
preferably are water or tetrahydrofuran, especially water and tetrahydrofuran as solvent. Dioxan and diethylene glycol dimethylether are also preferred solvents which may optionally (and preferably) be employed together with water. Preferably, the solvent will be present in an amount of between 3 and 10vol relative to the amount of the starting material (1wt.), more preferably between 4 and 6 vol., especially 5 vol. Preferably the oxidising agent is present in an amount of 1-9 molar equivalents relative to the amount of the starting material. For example, when a 50% w/w aqueous solution of periodic acid is employed, the oxidising agent may be present in an amount of between 1.1 and 10wt. relative to the amount of the starting material (1wt.), more preferably between 1.1 and 3wt., especially 1.3wt. Preferably, the oxidation step will comprise the use of a chemical oxidising agent. More preferably, the oxidising agent will be periodic acid or iodic acid or a salt thereof. Most preferably, the oxidising agent will be periodic acid or sodium periodate, especially periodic acid. Alternatively (or in addition), it will also be appreciated that the oxidation step may comprise any suitable oxidation reaction, eg one which utilises air and/or oxygen. When the oxidation reaction utilises air and/or oxygen, the solvent used in said reaction will preferably be methanol. Preferably, step (a) will involve incubating the reagents at room temperature or a little warmer, say around 25°C eg for 2 hours. The compound of formula (IV) may be isolated by recrystallisation from the reaction mixture by addition of an anti-solvent. A suitable anti-solvent for compound of formula (IV) is water. Surprisingly we have discovered that it is highly desirable to control the conditions under which the compound of formula (IV) is precipitated by addition of anti-solvent eg water. When the recrystallisation is performed using chilled water (eg water/ice mixture at a temperature of 0-5 °C) although better anti-solvent properties may be expected we have found that the crystalline product produced is very voluminous, resembles a soft gel and is very difficult to filter. Without being limited by theory we believe that this low density product contains a large amount of solvated solvent within the crystal lattice. By contrast when conditions of around 10 °C or higher are used (eg around ambient temperature) a granular product of a sand like consistency which is very easily filtered is produced. Under these conditions, crystallisation typically commences after around 1 hour and is typically completed within a few hours (eg 2 hours). Without being limited by theory we believe that this granular product contains little or no solvated solvent within the crystal lattice.

Step (b) will typically comprise the addition of a reagent suitable for converting a carboxylic acid to a carbothioic acid eg using hydrogen sulphide gas together with a suitable coupling agent eg carbonyldiimidazole (CDI) in the presence of a suitable solvent eg dimethylformamide.

An alternative process for preparing a compound of formula (II) comprises treating a compound of formula (X) with a reagent suitable for converting a carboxylic acid to a carbothioic acid eg using hydrogen sulphide gas together with a suitable coupling agent such as CDI in the presence of a suitable solvent eg DMF. Compounds of formula (X) may be prepared by methodology analogous to that described herein.

An alternative process for preparing a compound of formula (I) or a solvate thereof comprises reacting a compound of formula (VI) with a fluorine source.

Examples of suitable sources of fluorine include fluoride (eg sodium fluoride) or, more preferably, HF. The preferred reagent is aqueous HF. A solvent such as THF or DMF may be employed.

A compound of formula (VI) may be prepared by a process comprising

(a) alkylating a compound of formula (VII)
or a salt thereof;
(b) reacting a compound of formula (VIII)

with an epoxide forming reagent; or
(c) esterifying a compound of formula (IX)

[0091] In process (a), analogous conditions to those described above for the conversion of a compound of formula (II) to a compound of formula (I) may be employed. Typically compound of formula (VII) will be reacted with a compound of formula FCH$_2$L wherein L represents a leaving group (eg a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane.

[0092] Process (b) is preferably performed in two steps: (i) formation of a halohydrin especially a bromohydrin (eg by reaction with bromodan or equivalent reagent), followed by (ii) treatment with base such as sodium hydroxide so as to effect ring closure. The product of step (i) is a compound of formula (IXA) which is a novel intermediate that may be isolated, if desired:
wherein X represents halogen, especially Br.

[0093] In process (c), a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride in the presence of an organic base eg triethylamine. This reaction may be performed at elevated temperature eg around 60 °C or else at ambient temperature in the presence of an acylation catalyst eg dimethylamino pyridine (DMAP).

[0094] Compounds of formula (VII) may be prepared by a process comprising esterification of a compound of formula (XI)

[0095] Analogous conditions to those described above for the conversion of a compound of formula (III) to a compound of formula (II) may be employed. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride in the presence of an organic base eg triethylamine. Compound of formula (XI) is known (J Labelled Compd Radiopharm (1997) 39(7) 567-584).

[0096] A compound of formula (VIII) may be prepared by a process comprising

(a) alkylating a compound of formula (XII)

or a salt thereof; or

(b) esterifying a compound of formula (XIII)
In process (a), analogous conditions to those described above for the conversion of a compound of formula (II) to a compound of formula (I) may be employed. Typically compound of formula (XII) will be reacted with a compound of formula $\text{FCH}_2\text{L}$ wherein L represents a leaving group (e.g., a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane.

In process (b), analogous conditions to those employed above for the conversion of a compound of formula (IX) to a compound of formula (VI) may be employed. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide e.g., 2-furoyl chloride in the presence of an organic base e.g., triethylamine.

Compounds of formula (IX) and (XIII) may be prepared by alkylating the corresponding thioacids (XI) and (XIV) (defined below) using methodology analogous to that already described (e.g., by reaction with a compound of formula $\text{FCH}_2\text{L}$ wherein L represents a leaving group (e.g., a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane. The thioacid (XI) is a known compound (J Labelled Compd Radiopharm (1997) 39(7) 567-584).

Compound of formula (XII) may be prepared by a process comprising esterifying a compound of formula (XIV): or a salt thereof.

This process may be performed using methodology analogous to that already described. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide e.g., 2-furoyl chloride in the presence of an organic base e.g., triethylamine.

Compounds of formula (XIV) may be prepared from the corresponding carboxylic acid e.g., by a process analogous to that described above for the conversion of a compound of formula (IV) to a compound of formula (III). The aforesaid corresponding carboxylic acid is known (Upjohn, WO 90/15816).

A further alternative process for preparing a compound of formula (I) or a solvate thereof comprises deprotecting or unmasking a compound of formula (I) in which the 11-β-hydroxy group is protected or masked. A first such process comprises deprotecting a compound of formula (XV)
wherein P represents a hydroxy protecting group.


**[0104]** Examples of suitable hydroxy protecting groups P include groups selected from carbonate, alkyl (eg t-butyl or methoxymethyl), aralkyl (eg benzyl, p-nitrobenzyl, diphenylmethyl or triphenylmethyl), heterocyclic groups such as tetrahydropyranyl, acyl (eg acetyl or benzyl) and silyl groups such as trialkylsilyl (eg t-butyldimethylsilyl). The hydroxy protecting groups may be removed by conventional techniques. Thus, for example, carbonate may be removed by treatment with base and alkyl, silyl, acyl and heterocyclic groups may be removed by solvolysis eg by hydrolysis under acid or basic conditions. Aralkyl groups such as triphenylmethyl may similarly be removed by solvolysis eg by hydrolysis under acidic conditions. Aralkyl groups such as benzyl or p-nitrobenzyl may be cleaved by hydrogenolysis in the presence of a Noble metal catalyst such as palladium on charcoal. p-Nitrobenzyl may also be cleaved by photolysis.

**[0105]** The 11-β-hydroxy group may be masked as a carbonyl group. Thus a second such process comprises reduction of a compound of formula (XVI)

**[0106]** Reduction to the compound of formula (I) may be achieved eg by treatment with a hydride reducing agent such as borohydride eg sodium borohydride.

**[0107]** The 11-ketone (XVI) may also be masked. Examples of masked derivatives of compound of formula (XVI) include (i) ketal derivatives eg ketals formed by treatment of the compound of formula (XVI) with an alcohol eg methanol, ethanol or ethan-1,2-diol, (ii) dithioketal derivatives eg dithioketals formed by treatment of the compound of formula (XVI) with a thiol eg methanethiol, ethanethiol or ethan-1,2-dithiol, (iii) monothioketal derivatives eg monothioketals formed by treatment of the compound of formula (XVI) with eg 1-hydroxy-ethane-2-thiol, (iv) derivatives formed by treatment of the compound of formula (XVI) with an alcoholic eg ephedrine, (v) imines formed by treatment of the compound of formula (XVI) with amines, (vi) oximes formed by treatment of compounds of formula (XVI) with hydroxylamines. We claims such derivatives of compound of formula (XVI) as an aspect of the invention.

**[0108]** These masked derivatives may be converted back to the ketone by conventional means eg ketals, imines and oximes are converted to carbonyl by treatment with dilute acid and dithioketals are converted to the ketone by a variety of methods as described by P. C. Bulman Page et al (1989), Tetrahedron, 45, 7643-7677 and references therein.

**[0109]** Compounds of formula (XV) may be prepared by a process comprising

(a) alkylating a compound of formula (XVII)
or a salt thereof wherein P represents a hydroxy protecting group; or
(b) esterifying a compound of formula (XVIII)

[0110] In step (a), analogous conditions to those described above for the conversion of a compound of formula (II) to a compound of formula (I) may be employed. Typically compound of formula (XVII) will be reacted with a compound of formula FCH₂L wherein L represents a leaving group (e.g., a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane.

[0111] In step (b), analogous conditions to those employed above for the conversion of a compound of formula (IX) to a compound of formula (VI) may be employed. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide e.g., 2-furoyl chloride in the presence of an organic base e.g., triethylamine.

[0112] Compound of formula (XVIII) may be prepared by alkylating the corresponding thioacid using methodology analogous to that already described (e.g., by reaction with a compound of formula FCH₂L wherein L represents a leaving group (e.g., a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane. The corresponding thioacids are known compounds or may be prepared by standard methodology. Compound of formula (XVIII) may alternatively be prepared by protection of the corresponding hydroxy derivative.

[0113] Compound of formula (XVII) may be prepared by a process comprising esterifying a compound of formula (XIX)
or a salt thereof wherein P represents a hydroxy protecting group.

This process may be performed using methodology analogous to that already described for the conversion of compounds of formula (III) to (II). For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride in the presence of an organic base eg triethylamine.

Compounds of formula (XIX) may be prepared by protecting the corresponding hydroxy derivative (111), having first protected the thioacid which would then be deprotected.

Compounds of formula (XVI) may be prepared by a process comprising

(a) alkylating a compound of formula (XX)

or a salt thereof or a derivative wherein the 11-carbonyl group is masked; or

(b) esterifying a compound of formula (XXI)

or a derivative wherein the 11-carbonyl group is masked.

In step (a), analogous conditions to those described above for the conversion of a compound of formula (III) to a compound of formula (II) may be employed. Typically compound of formula (XX) will be reacted with a compound of formula FCH₂L wherein L represents a leaving group (eg a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane.

In step (b), analogous conditions to those employed above for the conversion of a compound of formula (IX) to a compound of formula (VI) may be employed. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride in the presence of an organic base eg triethylamine.

Compound of formula (XXI) or a derivative thereof wherein the 11-ketone group is masked may be prepared by alkylating the corresponding thioacid using methodology analogous to that already described (eg by reaction with a compound of formula FCH₂L wherein L represents a leaving group (eg a halogen atom, a mesyl or tosyl group or the like) for example, an appropriate fluoromethyl halide under standard conditions. Preferably, the fluoromethyl halide reagent is bromofluoromethane. The corresponding thioacids are known compounds or may be prepared from the corresponding carboxylic acids by methods analogous to those previously described.

Compound of formula (XX) may be prepared by a process comprising esterifying a compound of formula (XXII)
or a derivative thereof wherein the 11-ketone group is masked.

This process may be performed using methodology analogous to that already described. For example, a suitable reagent would be an activated derivative of 2-furoic acid such as an activated ester or preferably a 2-furoyl halide eg 2-furoyl chloride in the presence of an organic base eg triethylamine.

Compounds of formula (XXII) and derivatives thereof wherein the 11-ketone is masked may be prepared by oxidation of the corresponding hydroxy derivative (IV) followed by masking of the ketone and subsequent conversion of the carboxylic acid group to the thioacid (see eg conversion of compounds of formula (IV) to (III)).

A further alternative process for the preparation of compounds of formula (I) or a solvate thereof comprises reaction of a compound of formula (XXIII) wherein L represents a leaving group (eg halide other than fluoride such as chloride, iodide or a sulphonate ester such as mesylate, tosylate, triflate) with a fluorine source.

Preferably the fluorine source is fluoride ion eg KF. Further details for this conversion may be obtained by reference to G. H. Phillipps et al., (1994) Journal of Medicinal Chemistry, 37, 3717-3729 or J Labelled Compd Radiopharm (1997) 39(7) 567-584.

Compounds of formula (XXIII) may be prepared by methods analogous to those described herein. Corresponding novel intermediates of formula (VI), (VIII), (IX), (IXA), (XV) and (XVI) wherein the -CH2F moiety is replaced with a -CH2L moiety (wherein L represents a leaving group other than fluorine) are claimed as an aspect of the invention.

A further alternative process for the preparation of compounds of formula (I) or a solvate thereof comprises deprotection or unmasking of a derivative of a compound of formula (I) in which the 3-carbonyl group is protected or masked.

The 3-carbonyl group may be masked in a manner analogous to that described above in relation to masking of the 11-carbonyl position. Thus the 3-carbonyl may be masked eg as a ketal, monothioketal, dithioketal, derivative with an alcoholamine, oxime or imine. The carbonyl group may be recovered by conventional means eg ketals are converted to carbonyl by treatment with dilute acid and dithioketals are converted to the ketone by a variety of methods as described by P. C. Bulman Page et al (1989), Tetrahedron, 45, 7643-7677 and references therein.

Certain intermediate compounds are new and we provide these, together where appropriate with their salts and solvates, as an aspect of the invention.

As noted above, we provide as a particular aspect of the invention a process for preparing a compound of formula (I) in unsolvated form which comprises:

(a) Crystallising the compound of formula (I) in the presence of a non-solvating solvent such as ethanol, methanol,
water, ethyl acetate, toluene, methylisobutylketone or mixtures thereof; or
(b) Desolvating a compound of formula (I) in solvated form (eg in the form of a solvate with acetone, isopropanol, methyl ethyl ketone, DMF or tetrahydrofuran) eg by heating.

[0129] In step (b) the desolvation will generally be performed at a temperature exceeding 50 °C preferably at a temperature exceeding 100 °C. Generally heating will be performed under vacuum.

[0130] There is also provided a compound of formula (I) in unsolvated form obtainable by the aforementioned process.

[0131] There is also provided as a particular aspect of the invention a process for preparing a compound of formula (I) as unsolvated Form 1 polymorph which comprises dissolving compound of formula (I) in methyl isobutyl ketone, ethyl acetate or methyl acetate and producing compound of formula (I) as unsolvated Form 1 by addition of a non-solvating anti-solvent such as iso-octane or toluene.

[0132] According to a first preferred embodiment of this process the compound of formula (I) may be dissolved in ethyl acetate and compound of formula (I) as unsolvated Form 1 polymorph may be obtained by addition of toluene as anti-solvent. In order to improve the yield, preferably the ethyl acetate solution is hot and once the toluene has been added the mixture is distilled to reduce the content of ethyl acetate.

[0133] According to a second preferred embodiment of this process the compound of formula (I) may be dissolved in methyl isobutyl ketone and compound of formula (I) as unsolvated Form 1 polymorph may be obtained by addition of iso-octane as anti-solvent.

[0134] There is also provided a compound of formula (I) as unsolvated Form 1 polymorph obtainable by the aforementioned processes.

[0135] A process for preparing a compound of formula (I) as unsolvated Form 2 polymorph comprises dissolving compound of formula (I) in unsolvated form in methanol or dry dichloromethane and recrystallising the compound of formula (I) as unsolvated Form 2 polymorph. Typically the compound of formula (I) will be dissolved in hot in methanol or dry dichloromethane and allowed to cool.

[0136] There is also provided a compound of formula (I) as unsolvated Form 2 polymorph obtainable by the aforementioned process.

[0137] A process for preparing a compound of formula (I) as unsolvated Form 3 polymorph comprises dissolving compound of formula (I) or a solvate thereof (in particular as the acetone solvate) in dichloromethane in the presence of water (typically 1-3% water by volume) and recrystallising the compound of formula (I) as unsolvated Form 3 polymorph.

[0138] There is also provided a compound of formula (I) as unsolvated Form 3 polymorph obtainable by the aforementioned process.

[0139] The advantages of the compound of formula (I) and/or its solvates or polymorphs may include the fact that the substance appears to demonstrate excellent anti-inflammatory properties, with predictable pharmacokinetic and pharmacodynamic behaviour, with an attractive side-effect profile and is compatible with a convenient regime of treatment in human patients. Further advantages may include the fact that the substance has desirable physical and chemical properties which allow for ready manufacture and storage.

Brief Description of the Figures:

[0140] The following non-limiting Examples illustrate the invention:

Figure 1: Overlay of the XRPD profiles of Form 1, Form 2 and Form 3 polymorphs of unsolvated Compound of formula (I)
Figure 2: Overlay of the XRPD profiles of Form 1, Form 2 and a 50:50 mixture of Form 1 and Form 2 polymorphs of unsolvated Compound of formula (I) together with the time dependence of the profile of the 50:50 mixture of Form 1 and Form 2
Figure 3: DSC and TGA profiles of Form 1 polymorph of Unsolvated Compound of formula (I)
Figure 4: Temperature dependence of the XRPD profile of Compound of formula (I) Unsolvated Form 3 obtained at 5 timepoints
Figure 5: Temperature and time profile for the XRPD experiments of Figure 4
EXAMPLES

General

[0142] 1H-nmr spectra were recorded at 400 MHz and the chemical shifts are expressed in ppm relative to tetramethylsilane. The following abbreviations are used to describe the multiplicities of the signals: s (singlet), d (doublet), t (triplet), q (quartet), m (multiplet), dd (doublet of doublets), ddd (doublet of triplet of doublets), dt (doublet of triplets) and b (broad). Biotage refers to prepacked silica gel cartridges containing KP-Sil run on flash 12i chromatography module. LCMS was conducted on a Supelcosil LCABZ+PLUS column (3.3 cm x 4.6 mm ID) eluting with 0.1% HCO2H and 0.01 M ammonium acetate in water (solvent A), and 0.05% HCO2H 5% water in acetonitrile (solvent B), using the following elution gradient 0-0.7 min 0%B, 0.7-4.2 min 100%B, 4.2-5.3 min 0%B, 5.3-5.5 min 0%B at a flow rate of 3 ml/min. The mass spectra were recorded on a Fisons VG Platform spectrometer using electrospray positive and negative mode (ES+ve and ES-ve).

[0143] DSC and TGA profiles were obtained using a Netzsch STA449C simultaneous thermal analyser and using an unsealed pan with nitrogen gas flow and a thermal gradient of 10 °C/min. The moisture sorption characteristics were obtained using a Hiden Igasorb water sorption microbalance. The programme provides for stepwise increase in relative humidity (RH) from 0% to 90% RH and then decrease back to 0% RH in steps of 10% RH.

[0145] The XRPD analysis shown in Figure 1 and 2 were performed on a Phillips X’pert MPD powder diffractometer, serial number DY667. The method runs from 2 to 45 degrees 2Theta with 0.02 degree 2Theta step size and a 1 second collection time at each step. The XRPD analysis shown in Figure 4 employed the same instrument with an Anton Parr TTK thermal accessory using a method running from 2 to 35 degrees 2Theta with 0.04 degree 2Theta step size and a 1 second collection time.

Intermediates

Intermediate 1: 6α, 9α-Difluoro-17α-[[2-furanylcarbonyl]oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1, 4-diene-17β-carbothioic acid

[0146] A solution of 6α, 9α-difluoro-11β, 17α-dihydroxy-16α-methyl-3-oxo-androsta-1, 4-diene-17β-carbothioic acid (prepared in accordance with the procedure described in GB 2088877B) (18g, 43.64mmol) in anhydrous dichloromethane (200ml) and triethylamine (15.94ml, 114mmol) was treated at <5°C with a solution of 2-furoyl chloride (11.24ml, 114mmol) (prepared in accordance with the procedure described in GB 2088877B) (18g, 43.64mmol) in anhydrous dichloromethane (200ml) and triethylamine (15.94ml, 114mmol) in anhydrous dichloromethane (100ml) over approximately 40min. The solution was stirred at <5°C for 30min. The resulting solid was collected by filtration, washed successively with 3.5% sodium hydrogen carbonate solution, water, 1 M hydrochloric acid, and water and dried in vacuo at 60 °C to give a cream coloured solid. The dichloromethane filtrate was washed successively with 3.5% aqueous sodium hydroxide solution, water, 1 M hydrochloric acid, and water and dried in vacuo at 60 °C to give a cream coloured solid. The dichloromethane filtrate was washed successively with 3.5% sodium hydrogen carbonate solution, water, 1 M hydrochloric acid, water, dried (Na2SO4) and evaporated to give a cream coloured solid which was combined with that isolated above. The combined solids (26.9g) were suspended in acetone (450ml) and stirred. Diethylamine (16.8ml, 162mmol) was added and the mixture stirred at room temperature for 4.5h. The mixture was concentrated and the precipitate collected by filtration and washed with a little acetone. The washings and filtrate were combined, concentrated and loaded onto a silica gel Biotage column which was eluted with 24:1 chloroform: methanol. Fractions which contained the more polar component were combined and evaporated to give a cream coloured solid. The following abbreviations are used to describe the multiplicities of the signals: s (singlet), d (doublet), t (triplet), q (quartet), m (multiplet), dd (doublet of doublets), ddd (doublet of triplet of doublets), dt (doublet of triplets) and b (broad). Biotage refers to prepacked silica gel cartridges containing KP-Sil run on flash 12i chromatography module. LCMS was conducted on a Supelcosil LCABZ+PLUS column (3.3 cm x 4.6 mm ID) eluting with 0.1% HCO2H and 0.01 M ammonium acetate in water (solvent A), and 0.05% HCO2H 5% water in acetonitrile (solvent B), using the following elution gradient 0-0.7 min 0%B, 0.7-4.2 min 100%B, 4.2-5.3 min 0%B, 5.3-5.5 min 0%B at a flow rate of 3 ml/min. The mass spectra were recorded on a Fisons VG Platform spectrometer using electrospray positive and negative mode (ES+ve and ES-ve).

[0147] A stirred suspension of 6α, 9α-difluoro-11β, 17α-dihydroxy-16α-methyl-3-oxo-androsta-1, 4-diene-17β-carbothioic acid (prepared in accordance with the procedure described in GB 2088877B) (1wt, 49.5g) in acetone (10vol) is cooled to 0-5°C and treated with triethylamine (0.51wt, 2.1eq), keeping the temperature below 5°C, and stirred for 5 min at 0-5°C. 2-Furoyl chloride (0.65wt, 2.05eq) is then added over a minimum of 20min, maintaining a reaction temperature at 0-5°C. The reaction is stirred for 30min at 0-5°C then sampled for analysis by HPLC. A solution of diethanolamine (1.02wt, 4eq) in methanol (0.8vol) is added over ca 15min followed by a line wash of methanol (0.2vol) and the reaction...
stirred at 0-5°C for 1 h. The reaction is again sampled for analysis by HPLC then warmed to approximately 20°C and treated with water (1:1.1v). The mixture is then treated with a solution of HCl (SG1.18 (11.5M), 1vol) in water (10vol) over ca 20min maintaining a reaction temperature below 25°C. The suspension is stirred at 20-23°C for at least 30 minutes then filtered. The filter cake is washed with water (3x2vol). The product is dried in vacuo at approximately 60°C overnight to give the title compound as a white solid (58.7g, 96.5%).

Intermediate 1: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid (second alternative method)

[0148] A stirred suspension of 6α, 9α-difluoro-11β, 17α-dihydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid (prepared in accordance with the procedure described in GB 2088877B) (1wt, 49.5g) in acetone (10vol) is cooled to 0-5°C and treated with triethylamine (0.51wt, 2.1eq), keeping the temperature below 5°C, and stirred for 5 min at 0-5°. 2-Furoyl chloride (0.65wt, 2.05eq) is then added over a minimum of 20min, maintaining a reaction temperature at 0-5°C. The reaction mixture is stirred for at least 30 minutes and diluted with water (10vol) maintaining a reaction temperature in the range 0-5°C. The resultant precipitate is collected by filtration and washed sequentially with acetone/water (50/50 2vol) and water (2x2vol). The product is dried under vacuum at approximately 40-50°C (75.3g, 98.7%). NMR (CD3CN) 0.99 (3H, d) (J = 7.3Hz), 1.24 (3H, s), 1.38 (1H, m) (J = 3.9Hz), 1.54 (3H, s), 1.67 (1H, m), 1.89 (1H, broad d) (J = 15.2Hz), 1.9-2.0 (1H, m), 3.39 (1H, m), 4.33 (1H, m), 4.93 (1H, broad s), 5.53 (1H, ddd) (J = 6.9,1.9Hz, JHF = 50.9Hz), 6.24 (1H, m), 6.29 (1H, dd) (J = 10.3, 2.0Hz), 6.63 (2H, m), 7.24-7.31 (3H, m), 7.79 (1H, dd) (J = <1Hz), 7.86 (1H, dd) (J = <1Hz). A portion of the product (0.56g) is mixed with 6α, 9α-difluoro-11β, 17α-dihydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid (0.41g) in a 1:1 molar ratio in DMF (10volumes wrt total steroid input). The reaction mixture is treated with triethylamine (35ml) maintaining a reaction temperature in the range 0-5°C. The mixture is stirred at 0-5°C for 1 hour. A solution of diethanolamine (52.8g) in methanol (50ml) is added and the mixture is stirred at 0-5°C for 1 hour. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C. The organic phase is separated and the aqueous phase is treated with triethylamine (35ml) maintaining a reaction temperature in the range 0-5°C. The mixture is stirred for at least 2 hours. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C. The organic phase is separated and the aqueous phase is treated with triethylamine (35ml) maintaining a reaction temperature in the range 0-5°C. The mixture is stirred for at least 2 hours. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C. The organic phase is separated and the aqueous phase is treated with triethylamine (35ml) maintaining a reaction temperature in the range 0-5°C. The mixture is stirred for at least 2 hours. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C. The organic phase is separated and the aqueous phase is treated with triethylamine (35ml) maintaining a reaction temperature in the range 0-5°C. The mixture is stirred for at least 2 hours. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C.

Intermediate 1A: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-14-diene-17β-carbothioic acid disopropylethylamine salt

[0149] A stirred suspension of 6α, 9α-difluoro-11β, 17α-dihydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid (prepared in accordance with the procedure described in GB 2088877B) (49.5g) in methylacetate (500ml) is treated with triethylamine (52.8g) in methanol (50ml) and the mixture stirred at 0-5°C for 1 hour. A solution of diethanolamine (52.8g) in methanol (50ml) is added and the mixture stirred at 0-5°C for at least 2 hours. Dilute hydrochloric acid (approx 1M, 550ml) is added maintaining a reaction temperature below 15°C and the mixture stirred at 15°C. The organic phase is separated and the aqueous phase is back extracted with methyl acetate (2x250ml). All of the organic phases are combined, washed sequentially with brine (5 x 250ml) and treated with diisopropylethylamine (30ml). The reaction mixture is concentrated by distillation at atmospheric pressure to an approximate volume of 250ml and cooled to 25-30°C (crystallisation of the desired product normally occurs during distillation/subsequent cooling). Tertiary butyl methyl ether (TBME) (500ml) is added, the slurry further cooled and aged at 0-5°C for at least 10 minutes. The product is filtered off, washed with chilled TBME (2x200ml) and dried under vacuum at approximately 40-50°C (75.3g, 98.7%). NMR (CDCl3) δ: 7.54-7.48 (1H, m), 7.20-7.12 (1H, dd), 7.07-6.99 (1H, dd), 6.48-6.41 (2H, m), 6.41-6.32 (1H, dd), 5.51-5.28 (1H, dddd JHF 50Hz), 4.45-4.33 (1H, bd), 3.92-3.73 (3H, bm), 3.27-3.14 (2H, q), 2.64-2.12 (5H, m), 1.88-1.71 (2H, m), 1.58-1.15 (3H, s), 1.50-1.38 (15H, m), 1.32-1.23 (1H, m), 1.23-1.15 (3H s), 1.09-0.99 (3H, d).

Intermediate 1B: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid triethylamine salt

[0150] A stirred suspension of Intermediate 1 (30g) in ethylacetate (900ml) is treated with triethylamine (1.05 molar equivalents, 8.6ml) and the mixture is stirred at approximately 20°C for 1.5 hours. The precipitate is filtered off, washed with ethylacetate (2x2vol) and dried in vacuo at 45°C for 18 hours to give title compound as a white solid (28.8g, 80%). NMR (CDCl3) δ: 7.59-7.47 (1H, m), 7.23-7.13 (1H, dd), 7.08-6.99 (1H, d), 6.54-6.42 (2H, m), 6.42-6.32 (1H, dd), 5.55-5.26 (1H, dddd JHF 50Hz), 4.47-4.33 (1H, bd), 3.88-3.70 (1H, bm), 3.31-3.09 (6H, q), 2.66-2.14 (5H, m), 1.93-1.69 (2H, m), 1.61-1.48 (3H, s), 1.43-1.33 (9H, t), 1.33-1.26 (1H, m), 1.26-1.15 (3H s), 1.11-0.97 (3H, d).
Examples

Example 1: 6α, 9α-Difluoro-17α-[(2-furanyl carbonyl)oxy]-11β-hydroxy-16α-methyl-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester Unsolvated Form 1

[0151] A suspension of Intermediate 1 (2.5g, 4.94mmol) was dissolved in anhydrous N, N-dimethylformamide (25ml) and sodium hydrogen carbonate (465mg, 5.53mmol) was added. The mixture was stirred at -20°C and bromofluoromethane (0.77ml, 6.37mmol) was added and the mixture stirred at -20°C for 2h. Diethylamine (2.57ml, 24.7mmol) was added and the mixture stirred at -20°C for 30min. The mixture was added to 2M hydrochloric acid (93ml) and stirred for 30min. Water (300ml) was added and the precipitate was collected by filtration, washed with water and dried in vacuo at 50°C to give a white solid which was recrystallised from acetone/water (to yield the acetone solvate of 6α, 9α-difluoro-17α-[(2-furanyl carbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester) and dried in vacuo at 50°C to give the title compound (2.351g, 88%): LCMS retention time 3.66min, m/z 539 MH+

NMR δ (CDCl3) includes 7.60 (1H, m), 7.18 - 7.11 (2H, m), 6.52 (1H, dd, J 4.2Hz), 6.46 (1H, s), 6.41 (1H, dd, J 10, 2Hz), 5.95 and 5.82 (2H dd, J 51, 9Hz), 5.48 and 5.35 (1H, 2m), 4.48 (1H, m), 3.48 (1H, m), 1.55 (3H, s), 1.16 (3H, s), 1.06 (3H, d, J 7Hz).

Pharmacological Activity

In Vitro Pharmacological Activity

[0152] Pharmacological activity was assessed in a functional in vitro assay of glucocorticoid agonist activity which is generally predictive of anti-inflammatory or anti-allergic activity in vivo.

For the experiments in this section, compound of formula (I) was used as unsolvated Form 1. The functional assay was based on that described by K.P.Ray et al., Biochem J. (1997), 328, 707-715. A549 cells stably transfected with a reporter gene containing the NF-κB responsive elements from the ELAM gene promoter coupled to sPAP (secreted alkaline phosphatase) were treated with test compounds at appropriate doses for 1 hour at 37°C. The cells were then stimulated with tumour necrosis factor (TNF, 10ng/ml) for 16 hours, at which time the amount of alkaline phosphatase produced is measured by a standard colourimetric assay. Dose response curves were constructed from which EC50 values were estimated.

[0153] In this test the compound of Example 1 showed an EC50 value of <1nM.

[0154] The glucocorticoid receptor (GR) can function in at least two distinct mechanisms, by upregulating gene expression through the direct binding of GR to specific sequences in gene promoters, and by downregulating gene expression that is being driven by other transcription factors (such as NFκB or AP-1) through their direct interaction with GR.

[0155] In a variant of the above method, to monitor these functions, two reporter plasmids have been generated and introduced separately into A549 human lung epithelial cells by transfection. The first cell line contains the firefly luciferase reporter gene under the control of a synthetic promoter that specifically responds to activation of the transcription factor NFκB when stimulated with TNFα. The second cell line contains the renilla luciferase reporter gene under the control of a synthetic promoter that comprises 3 copies of the consensus glucocorticoid response element, and which responds to direct stimulation by glucocorticoids. Simultaneous measurement of transactivation and transrepression was conducted by mixing the two cell lines in a 1:1 ratio in 96 well plate (40,000 cells per well) and growing overnight at 37°C. Test compounds were dissolved in DMSO, and added to the cells at a final DMSO concentration of 0.7%. After incubation for 1h 0.5ng/ml TNFα (R&D Systems) was added and after a further 15 hours at 37°C, the levels of firefly and renilla luciferase were measured using the Packard Firelite kit following the manufacturers’ directions. Dose response curves were constructed from which EC50 values were determined.

<table>
<thead>
<tr>
<th>Transactivation (GR)</th>
<th>Transrepression (NFκB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ED50 (nM)</td>
<td>ED50 (nM)</td>
</tr>
<tr>
<td>Compound of Formula (I)</td>
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</tr>
<tr>
<td>Metabolite (X)</td>
<td>&gt;250</td>
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<tr>
<td>Fluticasone propionate</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>ED50 (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound of Formula (I)</td>
</tr>
<tr>
<td>Metabolite (X)</td>
</tr>
<tr>
<td>Fluticasone propionate</td>
</tr>
</tbody>
</table>

In Vivo Pharmacological Activity

[0156] Pharmacological activity in vivo was assessed in an ovalbumin sensitised Brown Norway rat eosinophilia model. This model is designed to mimic allergen induced lung eosinophilia, a major component of lung inflammation in asthma.

[0157] For the experiments in this section, compound of formula (I) was used as unsolvated Form 1.
Compound (1) produced dose dependent inhibition of lung eosinophilia in this model after dosing as an intra-tracheal (IT) suspension in saline 30 min prior to ovalbumin challenge. Significant inhibition is achieved after a single dose of 30µg of compound (I) and the response was significantly (p=0.016) greater than that seen with an equivalent dose of fluticasone propionate in the same study (69% inhibition with compound (I) vs 41% inhibition with fluticasone propionate).

In a rat model of thymus involution 3 daily IT doses of 100µg of compound (I) induced significantly smaller reductions in thymus weight (p=0.004) than an equivalent dose of fluticasone propionate in the same study. (67% reduction of thymus weight with compound (I) vs 78% reduction with fluticasone propionate).

Taken together these results indicate a superior therapeutic index for compound (I) compared to fluticasone propionate.

*In vitro* metabolism in rat and human hepatocytes

Incubation of compound (I) with rat or human hepatocytes shows the compound to be metabolised in an identical manner to fluticasone propionate with the 17-β carboxylic acid (X) being the only significant metabolite produced. Investigation of the rate of appearance of this metabolite on incubation of compound (I) with human hepatocytes (37°C, 10µM drug concentration, hepatocytes from 3 subjects, 0.2 and 0.7 million cells/mL) shows compound (I) to be metabolised ca. 5-fold more rapidly than fluticasone propionate:-

<table>
<thead>
<tr>
<th>Subject number</th>
<th>Cell density (million cells/mL)</th>
<th>17-β acid metabolite production (pmol/h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Compound (I)</td>
</tr>
<tr>
<td>1</td>
<td>0.2</td>
<td>48.9</td>
</tr>
<tr>
<td>1</td>
<td>0.7</td>
<td>73.3</td>
</tr>
<tr>
<td>2</td>
<td>0.2</td>
<td>118</td>
</tr>
<tr>
<td>2</td>
<td>0.7</td>
<td>903</td>
</tr>
<tr>
<td>3</td>
<td>0.2</td>
<td>102</td>
</tr>
<tr>
<td>3</td>
<td>0.7</td>
<td>580</td>
</tr>
</tbody>
</table>

Median metabolite production 102-118 pmol/h for compound (I) and 18.8-23.0 pmol/h for fluticasone propionate.

Pharmacokinetics after intravenous (IV) and oral dosing in rats

Compound (I) was dosed orally (0.1mg/kg) and IV (0.1 mg/kg) to male Wistar Han rats and pharmacokinetic parameters determined. Compound (I) showed negligible oral bioavailability (0.9%) and plasma clearance of 47.3 mL/min/kg, approaching liver blood flow (plasma clearance of fluticasone propionate = 45.2 mL/min/kg).

Pharmacokinetics after intra-tracheal dry powder dosing in the pig.

Anaesthetised pigs (2) were dosed intra-tracheally with a homogenous mixture of compound (I) (1mg) and fluticasone propionate (1mg) as a dry powder blend in lactose (10% w/w). Serial blood samples were taken for up to 8h following dosing. Plasma levels of compound (I) and fluticasone propionate were determined following extraction and analysis using LC-MS/MS methodology, the lower limits of quantitation of the methods were 10 and 20pg/mL for compound (I) and fluticasone propionate respectively. Using these methods compound (I) was quantifiable up to 2 hours after dosing and fluticasone propionate was quantifiable up to 8 hours after dosing. Maximum plasma concentrations were observed for both compounds within 15min after dosing. Plasma half-life data obtained from IV dosing (0.1mg/kg) was used to calculate AUC (0-inf) values for compound (I). This compensates for the plasma profile of Compound (I) only being defined up to 2 hours after an IT dose and removes any bias due to limited data between compound (I) and fluticasone propionate.

C<sub>max</sub> and AUC (0-inf) values show markedly reduced systemic exposure to compound (I) compared to fluticasone propionate:-
The pharmacokinetic parameters for both compound (I) and fluticasone propionate were the same in the anaesthetised pig following intravenous administration of a mixture of the two compounds at 0.1mg/kg. The clearance of these two glucocorticoids is similar is this experimental pig model.

Example 1: A mobile suspension of Intermediate 1A (12.61g, 19.8mmol; equivalent to 10g of Intermediate 1) in ethyl acetate (230ml) and water (50ml) is treated with a phase transfer catalyst (benzyltributylammonium chloride, 10mol%), cooled to 3°C and treated with bromofluoromethane (1.10ml, 19.5mmol, 0.98 equivalents), washing in with prechilled (0°C) (230ml) and water (50ml) is treated with a phase transfer catalyst (benzyltributylammonium chloride, 10mol%), cooled to 60°C to constant weight to yield the title compound (8.77g, 82%).

Example 2: A suspension of the title compound (150ml) and sucked dry. The product was dried at approximately 60°C under vacuum for 16h to leave the title compound (37.8g, 83.7%).

The XRPD pattern of Example 1 product is shown in Figure 1. The DSC and TGA profiles are shown in Figure 3.

Unsolvated Form 1

Example 2: A suspension of 6α, 9α-difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester acetone solvate (prepared eg according to Example 11) (50.0g) in acetone (1500ml) and water (75ml) was heated to reflux. The resultant mixture was clarified by hot filtration (Whatman 54 filter paper) and the solution was distilled at atmospheric pressure (approx 100ml solvent collected) giving crystallisation at a temperature of approximately 60°C under vacuum for 16h to leave the title compound (37.8g, 83.7%).

Example of Compound of Formula (I) Cmax (pg/mL) AUC (0-inf) (hr.pg/mL)
Pig 1 Pig 2 Pig 1 Pig 2
Fluticasone propionate 277 218 455 495

Example 1: 6α, 9α-Difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester Unsolvated Form 1 (first alternative method)

Example 2: 6α, 9α-Difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester Unsolvated Form 2

Example of Compound of Formula (I) Cmax (pg/mL) AUC (0-inf) (hr.pg/mL)
Pig 1 Pig 2 Pig 1 Pig 2
Fluticasone propionate 277 218 455 495

A more pure sample of 6α, 9α-difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1, first method) in methanol (60 volumes, distilled at atmospheric pressure) during which time some solid crystallised in the filtrate. Further acetone (200ml) was added to the filtrate giving a bright solution at reflux. The mixture was held at reflux for approximately 30 minutes and slowly cooled to ambient temperature. The mixture was further cooled and aged at 10-20°C for 2 hours. The slurry was cooled to below 10°C and the product was filtered off, sucked dry and dried at approximately 80°C under vacuum overnight to leave a white solid (4.34g, 71%).

A suspension of 6α, 9α-difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1, first method) in dichloromethane (180ml) was heated to reflux giving a bright solution. The solution was clarified by hot filtration (Whatman 54 filter paper) and the solution was distilled at atmospheric pressure (approx 100ml solvent collected) giving crystallisation at reflux. The mixture was held at reflux for approximately 30 minutes and slowly cooled to ambient temperature. The mixture was further cooled and aged at 10-20°C for 2 hours. The slurry was cooled to below 10°C and the product was filtered off, sucked dry and dried at approximately 80°C under vacuum overnight to leave a white solid (4.34g, 71%).

A suspension of 6α, 9α-difluoro-17α-[(2-furanyl(carbonyl)oxy)-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1, first method) in dichloromethane (180ml) was heated to reflux giving a bright solution. The solution was clarified by hot filtration (Whatman 54 filter paper) and the solution was distilled at atmospheric pressure (approx 100ml solvent collected) giving crystallisation at reflux. The mixture was held at reflux for approximately 30 minutes and slowly cooled to ambient temperature. The mixture was further cooled and aged at 10-20°C for 2 hours. The slurry was cooled to below 10°C and the product was filtered off, sucked dry and dried at approximately 80°C under vacuum overnight to leave a white solid (4.34g, 71%).
pressure to approx 37.5 volumes). The product was isolated by filtration and oven dried at 60°C under vacuum for 16 hours to leave a white, electrostatic solid (4.34g, 71%).

Example 2: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

Unsolvated Form 3

A suspension of 6α, 9α-difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester acetone solvate (prepared eg according to Example 11) (20.0g) in dichloromethane (800ml, 40 volumes) and water (10ml, 0.5 volumes) was heated to reflux giving a bright solution. The solution was clarified by hot filtration (Whatman 54 filter paper) during which time some solid crystallised in the filtrate which was fully dissolved upon heating to reflux. The solution was distilled at atmospheric pressure (approx 400ml solvent collected) and allowed to cool to ambient temperature. The mixture was further cooled and aged at <10°C for 10 minutes. The product was filtered off, sucked dry and dried at approximately 60°C under vacuum overnight to leave a white solid (12.7g, 70%).

Example 3: Interconversion of Forms 1, 2 and 3 of unsolvated 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

Slurrying a mixture of Form 1 and Form 2 in water at ambient temperature revealed that the components are transformed entirely to Form 1 with time. XRPD results are shown in Figure 2. Similar results were obtained by slurrying a mixture of Form 1 and Form 2 in ethanol at ambient temperature. From these results it may be concluded that Form 1 is the thermodynamically more stable polymorphic form out of the two forms.

Thermal XRPD studies on Form 3 were performed as shown in Figure 4. The temperature and time profile is shown in Figure 5 and the 5 traces shown in Figure 4 were obtained at the equilibration points shown in Figure 5. The results indicate that Form 3 is converted first to Form 2 and then to Form 1 as temperature is elevated.

Example 4: Moisture sorption of Forms 1, 2 and 3 of Unsolvated 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

The moisture sorption characteristics of the three forms were determined by monitoring the weight change of solid when exposed to stepwise increased and then decreased humidity. The results obtained were as follows:

Form 1: uptake of 0.18% w/w of moisture over the range 0-90% relative humidity at 25 °C.
Form 2: uptake of 1.1-2.4% w/w of moisture over the range 0-90% relative humidity at 25 °C.
Form 3: uptake of 1.2-2.5% w/w of moisture over the range 0-90% relative humidity at 25 °C.

Example 5: Enthalpy of dissolution of Forms 1 and 3 of unsolvated 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

Enthalpies of dissolution in DMSO and acetonitrile were determined at 25 °C. The results were as follows:

<table>
<thead>
<tr>
<th></th>
<th>Form I</th>
<th>Form 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetonitrile</td>
<td>+13.74</td>
<td>+8.62</td>
</tr>
<tr>
<td>DMSO</td>
<td>+1.46</td>
<td>-5.21</td>
</tr>
</tbody>
</table>

(results in kJ/mol)

Form these results it may be determined that the enthalpy of transition from Form 3 to Form 1 is approximately 5.1-6.7 kJ/mol. On the assumption that the entropy of transition is small, since both Forms are unsolvated, the enthalpy of transition may be equated with the free energy of transition. Thus these data suggest that Form 1 is the thermodynamically most stable form at 25 °C.
Example 7: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

**Methylethylketone solvate**

[0180] A suspension of 6α, 9α-difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1) (400mg) in methylethylketone (3.2ml) is heated to reflux giving a clear solution. A portion of the solvent is distilled off at atmospheric pressure (approx 1ml) and the mixture cooled to approximately 20°C. The crystallised product is filtered off, dried at approximately 20°C under vacuum to leave the title compound as a white solid (310mg, 68%). NMR \( \delta \) (CDCl₃) includes the peaks described in Example 1 for the parent compound and the following additional solvent peaks: 2.45 (2H, q), 2.14 (3H, s), 1.06 (3H, t).

Example 8: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

**Isopropanol solvate**

[0181] A solution of 6α, 9α-difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1) (150mg) in isopropanol (15ml) is left to slowly crystallise over a period of approximately 8 weeks. The resultant chunky crystals are isolated by filtration to leave the title compound as a white solid. NMR \( \delta \) (CDCl₃) includes the peaks described in Example 1 for the parent compound and the following additional solvent peaks: 4.03 (1H, m), 1.20 (6H, d).

Example 9: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

**Tetrahydrofuran solvate**

[0182] A suspension of 6α, 9α-difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester (prepared eg according to Example 1) (150mg) in THF (20vol) is warmed to give a clear solution. The solvent is allowed to slowly evaporate over a period of approximately 8 hours. The resultant chunky crystals are isolated by filtration to leave the title compound as a white solid. NMR \( \delta \) (CDCl₃) includes the peaks described in Example 1 for the parent compound and the following additional solvent peaks: 3.74 (4H, m), 1.85 (4H, m).

Example 9: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

**Tetrahydrofuran solvate (alternative method)**

[0183] A mobile suspension of 6α, 9α-difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid triethylamine salt (prepared eg according to Intermediate 1B) (1.2g) in THF (10ml) is treated with a phase transfer catalyst (tetrabutylammonium bromide, typically between 8 and 14mol%), cooled to approximately 3°C and treated with bromofluoromethane (0.98 equivalents). The suspension is stirred for between 2 and 5 hours, allowing to warm to 17°C. The reaction mixture is poured into water (30vol), stirred at approximately 10°C for 30 minutes and filtered off. The collected solid is washed with water (4x3vol) and the product oven dried under vacuum at 60°C overnight to give the title compound as a white solid (0.85g, 87%).

Example 10: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

**DMF solvate**

[0184] A mixture of Intermediate 1 (4.5g, 8.88 mmol) in DMF (31 ml) is treated with potassium bicarbonate (0.89g, 8.88mmol) and the mixture is cooled to -20°C. A solution of bromofluoromethane (0.95g, 8.50 mmol, 0.98 eqv.) in DMF (4.8 ml) at 0°C is added and the mixture is stirred at -20°C for 4 hours. The mixture is then stirred at -20°C for a further 30 minutes, added to 2M hydrochloric acid (100ml) and stirred for a further 30 minutes at 0-5 °C. The precipitate collected
by vacuum filtration, washed with water and dried at 50°C to give the title compound (4.47g, 82%). NMR δ (CD$_3$OD) includes the peaks described in Example 1 for the parent compound and the following additional solvent peaks: 7.98 (1H, bs), 2.99 (3H, s), 2.86 (3H, s).

Example 11: 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester

Acetone solvate

A solution of Intermediate 1 (530.1g, 1wt) in dimethylformamide (DMF) (8vol) is treated with potassium hydrogen carbonate (0.202wt, 1.02eq) and the mixture cooled to -17°C with stirring. Bromofluoromethane (BFM) (0.22wt, 0.99eq) is then added and the reaction stirred at -17°C for at least 2h. The reaction mixture is then added to water (17vol) at 5°C over ca 10min followed by a water (1vol) line wash. The suspension is stirred at 5-10°C for at least 30min and then filtered. The filter cake (the DMF solvate of 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester) is washed with water (4x4vol) and the product is pulled dry on the filter. The damp cake is returned to the vessel, acetone (5.75vol) added and heated at reflux for 2h. The mixture is cooled to 52°C and water (5.75vol) added, keeping temperature at 52°C. The mixture is then cooled to 20°C, filtered and dried in vacuo at 60°C overnight to give the title compound as a white solid (556.5g, 89%). NMR δ (CDCl$_3$) includes the peaks described in Example 1 for the parent compound and the following additional solvent peaks: 2.17 (6H, s).

Example 12: Dry powder composition containing 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, Unsolvated Form 1

[0185] A dry powder formulation was prepared as follows:

6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, unsolvated Form 1 (prepared according to Example 1, first alternative method and micronised to a MMD of 3μm): 0.20mg

Milled lactose (wherein not greater than 85% of particles have a MMD of 60-90μm, and not less than 15% of particles have a MMD of less than 15μm): 12mg

Example 13: Aerosol formulation containing 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, Unsolvated Form 1

[0186] An aluminium canister was filled with a formulation as follows:

6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, Unsolvated Form 1 (prepared according to Example 1, first alternative method and micronised to a MMD of 3μm): 250μg

1,1,1,2-tetrafluoroethane: (amounts per actuation) to 50μl

Polysorbate 20 0.8mg

Example 14: Nasal formulation containing 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, Unsolvated Form 1

[0187] in a total amount suitable for 120 actuations and the canister was fitted with a metering valve adapted to dispense 50μl per actuation.

Example 15: Nasal formulation containing 6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester, Unsolvated Form 1

[0188] A formulation for intranasal delivery was prepared as follows:

6α, 9α-Difluoro-17α-[(2-furanylcarbonyl)oxy]-11β-hydroxy-16α-methyl-3-oxo-androsta-1,4-diene-17β-carbothioic acid S-fluoromethyl ester Unsolvated Form 1 (prepared according to Example 1, first alternative method, micronised): 10mg

Polysorbate 20 0.8mg
The formulation was fitted into a spraypump capable of delivering a plurality of metered doses (Valois).

Claims

1. A compound of formula (I)

and solvates thereof.

2. A compound of formula (I) as defined in claim 1 in unsolvated form.

3. A compound of formula (I) in unsolvated form as defined in claim 2 in the form of Form 1 polymorph, which has an XRPD profile with a peak at around 18.9 degrees 2 Theta.

4. A compound of formula (I) in unsolvated form as defined in claim 2 in the form of Form 2 polymorph, which has an XRPD profile with a peak at around 18.4 and 21.5 degrees 2 Theta.

5. A compound of formula (I) in unsolvated form as defined in claim 2 in the form of Form 3 polymorph which has an XRPD profiled with a peak at around 18.6 and 19.2 degrees 2 Theta.

6. A compound of formula (I) as defined in claim 1 as a crystalline solid in the form of an essentially stoichiometric solvate with acetone.

7. A compound of formula (I) as defined in claim 1 as a crystalline solid in the form of an essentially stoichiometric solvate with tetrahydrofuran.

8. A compound of formula (I) as defined in claim 1 as a crystalline solid in the form of an essentially stoichiometric solvate with isopropanol.

9. A compound of formula (I) as defined in claim 1 as a crystalline solid in the form of an essentially stoichiometric solvate with methylethylketone.

10. A compound of formula (I) as defined in claim 1 as a crystalline solid in the form of an essentially stoichiometric solvate with dimethylformamide.

11. A compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 for use in veterinary or human medicine.
12. Use of a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 for the manufacture of a medicament for the treatment of inflammatory and/or allergic conditions.

13. Use of a compound of formula (I) or a physiologically acceptable solvate thereof according to claim 12 for the treatment of an inflammatory condition of the nose.

14. Use as claimed in claim 13 wherein the inflammatory condition of the nose is rhinitis.

15. Use of a compound of formula (I) or a physiologically acceptable solvate thereof according to claim 12 for the treatment of conjunctivitis.

16. Use as claimed in any one of claims 13 to 15 wherein the compound of formula (I) or a physiologically acceptable solvate thereof is for administration by inhalation.

17. Use as claimed in any one of claims 13 to 16 wherein the compound of formula (I) or a physiologically acceptable solvate thereof is for administration locally.

18. Use as claimed in any one of claims 13 to 16 wherein the compound of formula (I) or a physiologically acceptable solvate thereof is for administration topically.

19. A pharmaceutical composition comprising a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 in admixture with one or more physiologically acceptable diluents or carriers.

20. A pharmaceutical composition as claimed in claim 19 comprising a compound of formula (I) as defined in claim 2.

21. A pharmaceutical composition as claimed in claim 19 comprising a compound of formula (I) as defined in claim 3.

22. A pharmaceutical composition as claimed in claim 19 comprising a compound of formula (I) as defined in claim 4.

23. A pharmaceutical composition as claimed in claim 19 comprising a compound of formula (I) as defined in claim 5.

24. A pharmaceutical composition according to any one of claims 19 to 23 wherein the composition is a spray.

25. A pharmaceutical composition according to any one of claims 19 to 23, wherein said compound of formula (I) or a physiologically acceptable solvate thereof is present in the amount of 0.001 to 10% by weight of said composition.

26. A pharmaceutical formulation comprising a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 in admixture with one or more physiologically acceptable diluents or carriers which is non-pressurised and adapted to be administered topically to the nasal cavity.

27. A pharmaceutical formulation according to claim 26 which contains water as the diluent or carrier.

28. A pharmaceutical aerosol formulation comprising a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 and a fluorocarbon or hydrogen-containing chlorofluorocarbon as propellant, optionally in combination with a surfactant and or a cosolvent.

29. A pharmaceutical aerosol formulation according to claim 28 which comprises a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 3, and a fluorocarbon or hydrogen-containing chlorofluorocarbon as propellant and a suspending agent which is soluble in the propellant.

30. A pharmaceutical aerosol formulation according to claim 29 wherein the suspending agent is an oligolactic acid or a derivative thereof.

31. A pharmaceutical aerosol formulation according to any one of claims 28 to 30 wherein the propellant is selected from 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoro-n-propane and mixtures thereof.

32. A pharmaceutical aerosol formulation according to claim 28 which consists essentially of a compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 3 optionally in combination with
another therapeutically active agent and a propellant selected from 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoro-n-propane and mixtures thereof.

33. A pharmaceutical composition according to any one of claims 19 to 32 which further comprises another therapeutically active agent.

34. A pharmaceutical composition according to claim 33 wherein said another therapeutically active agent is an anti-histamine, anti-inflammatory agent or antinfective agent.

35. A pharmaceutical composition according to claim 34 wherein said anti-histamine is methapyrilene or loratadine, said anti-inflammatory agent is an NSAID and said antinfective agent is an antibiotic or antiviral.

36. A pharmaceutical composition according to claim 33 in which said another therapeutically active agent is a PDE4 inhibitor.

37. A pharmaceutical composition according to claim 36, wherein the PDE4 inhibitor is at least one selected from the group consisting of (R)-(++)-1-(4-bromobenzyl)-4-[(3-cyclopentyloxy)-4-methoxyphenyl]-2-pyrrolidone; 3-(cyclopentyloxy-4-methoxyphenyl)-1-(4-N'-[N2-cyano-S-methyl-isothioureido]benzyl)-2-pyrrolidone; cis 4-cyano-4-(3-cyclopentyloxy-4-methoxyphenyl)cyclohexan-1-carboxylic acid; cis-[4-cyano-4-(3-cyclopropylmethoxy-4-difluoromethoxyphenyl)cyclohexan-1-ol]; (R)-(++)-ethyl [4-(3-cyclopentyloxy-4-methoxyphenyl)pyrrolidine-2-ylidene]acetate; (S)-(++)-ethyl [4-(3-cyclopentyloxy-4-methoxyphenyl)pyrrolidine-2-ylidene]acetate; and 2-carbomethoxy-4-cyano-4-(3-cyclopropylmethoxy-4-difluoromethoxyphenyl)cyclohexan-1-one.

38. Compound of formula (I) or a physiologically acceptable solvate thereof as defined in any one of claims 1 to 5 for use in the treatment of an inflammatory and/or an allergic condition.

39. A process for preparing a compound of formula (I) according to claim 1 or a solvate thereof which comprises alkylation of the 17β-carbothioic group of a compound of formula (II)

40. A process according to claim 39 wherein alkylation is performed by reacting the compound of formula (II) or a salt thereof with a fluoromethyl halide.

41. A process for preparing a compound of formula (I) as unsolvated Form 1 polymorph according to claim 3 which comprises:

(a) Crystallising the compound of formula (I) in the presence of a non-solvating solvent; or
(b) Desolvating a compound of formula (I) in solvated form.

42. A process for preparing a compound of formula (I) as unsolvated Form 1 polymorph according to claim 3 which comprises dissolving compound of formula (I) in methylisobutylketone, ethyl acetate or methyl acetate and producing compound of formula (I) as unsolvated Form 1 by addition of a non-solvating anti-solvent.

43. A compound of formula (II)
44. A compound of formula (II) as defined in claim 43 in the form of a solid crystalline salt.

45. A compound of formula (II) according to claim 44 in the form of the diisopropylethylamine salt.

46. A process for preparing a compound of formula (II) as defined in claim 43 which comprises:

(a) reacting a compound of formula (III)

with an activated derivative of 2-furoic acid in an amount of at least 2 moles of the activated derivative per mole of compound of formula (III) to yield a compound of formula (IIA)

and

(b) removal of the sulphur-linked 2-furoyl moiety from compound of formula (IIA) by reaction of the product of step (a) with an organic primary or secondary amine base capable of forming a water soluble 2-furoyl amide.

47. A process for preparing a compound of formula (II) as claimed in claim 46 which further comprises the steps of

(c1) when the product of step (b) is dissolved in a substantially water immiscible organic solvent, purifying the compound of formula (II) by washing out the amide by-product from step (b) with an aqueous wash, or

(c2) when the product of step (b) is dissolved in a water miscible solvent, purifying the compound of formula (II) by treating the product of step (b) with an aqueous medium so as to precipitate out pure compound of formula
48. A process for preparing a compound of formula (II) as defined in claim 43 which comprises:

(a) reacting a compound of formula (III) as defined in claim 46 with an activated derivative of 2-furoic acid in an amount of at least 2 moles of activated derivative per mole of compound of formula (III) to yield a compound of formula (IIA) as defined in claim 46; and

(b) removal of the sulphur-linked 2-furoyl moiety from compound of formula (IIA) by reaction of the product of step (a) with a further mole of compound of formula (III) to give two moles of compound of formula (II).

49. A compound of formula (IIA)

50. A compound of formula (VI)

51. A compound of formula (VII)

or a salt thereof.

52. A compound of formula (VIII)
53. A compound of formula (IXA) wherein X represents halogen.

54. A compound of formula (IXA) as claimed in claim 53. wherein X represents Br.

55. A compound of formula (XII) or a salt thereof.
56. A compound of formula (XV)

\[
\begin{align*}
\text{EP1 305 329 B2} \\
\text{35} \\
\text{5} \\
\text{10} \\
\text{15} \\
\text{20} \\
\text{25} \\
\text{30} \\
\text{35} \\
\text{40} \\
\text{45} \\
\text{50} \\
\text{55}
\end{align*}
\]

wherein P represents a hydroxy protecting group.

57. A compound of formula (XVI)

58. A compound of formula (XVII)

or a salt thereof wherein P represents a hydroxy protecting group.

59. A compound of formula (XX)
60. A compound of formula (XXIII)

![Chemical Structure](image)

wherein L represents a leaving group other than fluorine.

61. A process for preparing compound of formula (I) in unsolvated Form 2 polymorph as claimed in claim 4 which comprises dissolving compound of formula (I) in unsolvated form in methanol or dry dichloromethane and recrystallising the compound of formula (I) as unsolvated Form 2 polymorph.

62. A process for preparing compound of formula (I) in unsolvated Form 3 polymorph as claimed in claim 5 which comprises dissolving compound of formula (I) or a solvate thereof in dichloromethane in the presence of water and recrystallising the compound of formula (I) as unsolvated Form 3 polymorph.

63. A process for preparing a compound of formula (I) as defined in claim 1 or a solvate thereof which comprises reacting a compound of formula (VI)

![Chemical Structure](image)

with a fluorine source.

64. A process for preparing a compound of formula (I) or a solvate thereof as defined in claim 1 or a solvate thereof which comprises providing a compound of formula (I) in which the 11-β-hydroxy group is protected or masked, deprotecting or unmasking the compound to yield the compound of formula (I) or a solvate thereof.

65. A process according to claim 64 wherein the 11-β-hydroxy group is protected which comprises deprotecting a compound of formula (XV)
wherein P represents a hydroxy protecting group.

66. A process according to claim 64 wherein the 11-β-hydroxy group is masked which comprise reduction of a compound of formula (XVI)

or a derivative wherein the 11-carbonyl group is masked.

67. A process for the preparation of a compound of formula (I) as defined in claim 1 or a solvate thereof which comprises reaction of a compound of formula (XXIII)

wherein L represents a leaving group with a fluorine source.

68. A process for the preparation of a compound of formula (I) as defined in claim 1 or a solvate thereof which comprises deprotection or unmasking of a derivative of a compound of formula (I) in which the 3-carbonyl group is protected or masked.

69. A compound of formula (X)
70. A process for preparing a compound of formula (II) as defined in claim 43 which comprises treating a compound of formula (X) as defined in claim 69 with a reagent suitable for converting a carboxylic acid to a carbothioic acid.

71. A process for preparing a compound of formula (VI) as defined in claim 50 which comprises esterifying a compound of formula (IX) with an activated derivative of 2-furoic acid.

72. A process for preparing a compound of formula (VIII) as defined in claim 52 which comprises esterifying a compound of formula (XIII) with an activated derivative of 2-furoic acid.

73. A process for preparing a compound of formula (XII) as defined in claim 55 which comprises esterifying a compound of formula (XIV)
or a salt thereof, with an activated derivative of 2-furoic acid.

74. A process for preparing a compound of formula (XVI) as defined in claim 57 which comprises esterifying a compound of formula (XXI)

![Chemical Structure](image)

or a derivative wherein the 11-carbonyl group is masked, with an activated derivative of 2-furoic acid.

75. A process for preparing a compound of formula (XX) as defined in claim 59 which comprises esterifying a compound of formula (XXII)

![Chemical Structure](image)

or a derivative wherein the 11-ketone group is masked, with an activated derivative of 2-furoic acid.

76. A metered dose inhaler comprising a pharmaceutical aerosol formulation according to claim 28.

77. A metered dose inhaler as claimed in claim 76 wherein said pharmaceutical aerosol formulation is retained in a pressurized canister closed with a valve.

78. A metered dose inhaler of claims 76 or 77 wherein said pharmaceutical aerosol formulation has a particle size in the range of 1-10μm.

**Patentansprüche**

1. Verbindung der Formel (I):
und Solvate davon.

2. Verbindung der Formel (I) gemäß Anspruch 1 in unsolvatisierter Form.

3. Verbindung der Formel (I) in unsolvatisierter Form gemäß Anspruch 2 in der Form des Form-1-Polymorphs, das ein XRPD-Profil mit einem Peak bei ungefähr 18,9° 2θ hat.

4. Verbindung der Formel (I) in unsolvatisierter Form gemäß Anspruch 2 in der Form des Form-2-Polymorphs, das ein XRPD-Profil mit einem Peak bei ungefähr 18,4 und 21,5° 2θ hat.

5. Verbindung der Formel (I) in unsolvatisierter Form gemäß Anspruch 2 in der Form des Form-3-Polymorphs, das ein XRPD-Profil mit einem Peak bei ungefähr 18,6 und 19,2° 2θ hat.

6. Verbindung der Formel (I) gemäß Anspruch 1 als kristalliner Feststoff in der Form eines im wesentlichen stöchiometrischen Solvats mit Aceton.

7. Verbindung der Formel (I) gemäß Anspruch 1 als kristalliner Feststoff in der Form eines im wesentlichen stöchiometrischen Solvats mit Tetrahydrofuran.

8. Verbindung der Formel (I) gemäß Anspruch 1 als kristalliner Feststoff in der Form eines im wesentlichen stöchiometrischen Solvats mit Isopropanol.

9. Verbindung der Formel (I) gemäß Anspruch 1 als kristalliner Feststoff in der Form eines im wesentlichen stöchiometrischen Solvats mit Methyläthyketon.

10. Verbindung der Formel (I) gemäß Anspruch 1 als kristalliner Feststoff in der Form eines im wesentlichen stöchiometrischen Solvats mit Dimethylformamid.

11. Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 5 zur Verwendung in der Veterinär- oder Humanmedizin.

12. Verwendung einer Verbindung der Formel (I) oder eines physiologisch annehmbaren Solvats davon gemäß einem der Ansprüche 1 bis 5 zur Herstellung eines Medikaments zur Behandlung von entzündlichen und/oder allergischen Zuständen.

13. Verwendung einer Verbindung der Formel (I) oder eines physiologisch annehmbaren Solvats davon gemäß Anspruch 12 zur Behandlung eines Entzündungszustands der Nase.


15. Verwendung einer Verbindung der Formel (I) oder eines physiologisch annehmbaren Solvats davon Anspruch 12 zur Behandlung von Konjunktivitis.
16. Verwendung gemäß einem der Ansprüche 13 bis 15, worin die Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon zur Verabreichung durch Inhalation ist.

17. Verwendung gemäß einem der Ansprüche 13 bis 16, worin die Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon zur lokalen Verabreichung ist.

18. Verwendung gemäß einem der Ansprüche 13 bis 16, worin die Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon zur topischen Verabreichung ist.

19. Pharmazeutische Zusammensetzung, die eine Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 5 im Gemisch mit einem oder mehreren physiologisch annehmbaren Verdünnungsmitteln oder Trägern umfaßt.

20. Pharmazeutische Zusammensetzung gemäß Anspruch 19, die eine Verbindung der Formel (I) gemäß Anspruch 2 umfaßt.

21. Pharmazeutische Zusammensetzung gemäß Anspruch 19, die eine Verbindung der Formel (I) gemäß Anspruch 3 umfaßt.

22. Pharmazeutische Zusammensetzung gemäß Anspruch 19, die eine Verbindung der Formel (I) gemäß Anspruch 4 umfaßt.

23. Pharmazeutische Zusammensetzung gemäß Anspruch 19, die eine Verbindung der Formel (I) gemäß Anspruch 5 umfaßt.


25. Pharmazeutische Zusammensetzung gemäß einem der Ansprüche 19 bis 23, worin die Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon in der Menge von 0,001 bis 10 Gew.% der Zusammensetzung vorliegt.

26. Pharmazeutische Formulierung, die eine Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 5 in Mischung mit einem oder mehreren physiologisch annehmbaren Verdünnungsmitteln oder Trägern umfaßt, nicht druckverdichtet ist und zur topischen Verabreichung in der Nasenhöhle angepaßt ist.

27. Pharmazeutische Formulierung gemäß Anspruch 26, die Wasser als Verdünnungsmittel oder Träger enthält.

28. Pharmazeutische Aerosolformulierung, die eine Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 5 und einen Fluorkohlenstoff oder wasserstoffhaltigen Chlorkohlenstoff als Treibmittel umfaßt, gegebenenfalls in Kombination mit einem Tensid und/oder einem Kosolvens.

29. Pharmazeutische Aerosolformulierung gemäß Anspruch 28, die eine Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 3 und einen Fluorkohlenstoff oder wasserstoffhaltigen Chlorkohlenstoff als Treibmittel und ein Suspendiermittel, das im Treibmittel löslich ist, umfaßt.

30. Pharmazeutische Aerosolformulierung gemäß Anspruch 29, worin das Suspendiermittel eine Oligomilchsäure oder ein Derivat davon ist.

31. Pharmazeutische Aerosolformulierung gemäß einem der Ansprüche 28 bis 30, worin das Treibmittel ausgewählt ist aus 1,1,1,2-Tetrafluorethan, 1,1,1,2,3,3,3-Heptafluor-n-propan und Mischungen daraus.

32. Pharmazeutische Aerosolformulierung gemäß Anspruch 28, die im wesentlichen aus einer Verbindung der Formel (I) oder einem physiologisch annehmbaren Solvat davon gemäß einem der Ansprüche 1 bis 3, gegebenenfalls in Kombination mit einem weiteren Therapeutikum und einem Treibmittel besteht, das aus 1,1,1,2-Tetrafluorethan, 1,1,1,2,3,3,3-Heptafluor-n-propan und Mischungen daraus ausgewählt ist.
33. Pharmazeutische Zusammensetzung gemäß einem der Ansprüche 19 bis 32, die ferner ein weiteres Therapeutikum umfaßt.

34. Pharmazeutische Zusammensetzung gemäß Anspruch 33, worin das weitere Therapeutikum ein Antihistamin, ein entzündungshemmendes Mittel oder ein infektionshemmendes Mittel ist.

35. Pharmazeutische Zusammensetzung gemäß Anspruch 34, worin das Antihistamin Methapyrilen oder Loratadin ist, das entzündungshemmende Mittel ein NSAID ist und das infektionshemmende Mittel ein Antibiotikum oder ein antivirales Mittel ist.

36. Pharmazeutische Zusammensetzung gemäß Anspruch 33, in der das weitere Therapeutikum ein PDE4-Inhibitor ist.

37. Pharmazeutische Zusammensetzung gemäß Anspruch 36, worin der PDE4-Inhibitor mindestens einer ist, der aus der Gruppe ausgewählt ist, die aus (R)-(+)1-(4-Brombenzyl)-4-[(3-cycloptyloxy)-4-methoxyphenyl]-2-pyrrolidon, 3-(Cycloptyloxy)-4-methoxyphenyl]-1-(4-N'-[N2-cyano-S-methyl-isothioureido]benzyl)-2-pyrrolidon, cis-4-Cyano-4-(3-cycloptyloxy-4-methoxyphenyl)cyclohexan-1-carbonsäure, cis-4-(3-cycloptyloxy-4-difluormethoxyphenyl)cyclohexan-1-carbonsäure und 2-Carbomethoxy-4-cyano-4-(3-cyclopropylmethoxy-4-difluormethoxyphenyl)cyclohexan-1-on besteht.

38. Verbindung der Formel (I) oder ein physiologisch annehmbares Solvat davon gemäß einem der Ansprüche 1 bis 5 zur Verwendung in der Behandlung eines entzündlichen und/oder eines allergischen Zustands.

39. Verfahren zur Herstellung einer Verbindung der Formel (I) gemäß Anspruch 1 oder eines Solvats davon, das die Alkylierung der 17β-Thiocarbonsäuregruppe der Verbindung der Formel (II):

![Chemical Structure](image)

(II)

oder eines Salzes davon mit einer Verbindung der Formel FCH₃-L umfaßt, worin L eine Abgangsgruppe darstellt.

40. Verfahren gemäß Anspruch 39, worin die Alkylierung durchgeführt wird durch das Umsetzen der Verbindung der Formel (II) oder eines Salzes davon mit einem Fluormethylhalogenid.

41. Verfahren zur Herstellung einer Verbindung der Formel (I) als unsolvatisiertes Form-1-Polymorph gemäß Anspruch 3, umfassend:

(a) Kristallisieren der Verbindung der Formel (I) in Gegenwart eines nichtsolvatisierenden Lösungsmittels; oder

(b) Desolvatisieren einer Verbindung der Formel (I) in solvatisierter Form.

42. Verfahren zur Herstellung einer Verbindung der Formel (I) als unsolvatisiertes Form-1-Polymorph gemäß Anspruch 3, das das Lösen einer Verbindung der Formel (I) in Methylisobutylketon, Ethylacetat oder Methylacetat und das Herstellen einer Verbindung der Formel (I) als unsolvatisierte Form 1 durch Zugabe eines nichtsolvatisierenden Antilösungsmittels umfaßt.

43. Verbindung der Formel (II):
oder ein Salz davon.

44. Verbindung der Formel (II) gemäß Anspruch 43 in der Form eines festen kristallinen Salzes.

45. Verbindung der Formel (II) gemäß Anspruch 44 in der Form des Diisopropylethylaminsalzes.

46. Verfahren zur Herstellung einer Verbindung der Formel (II) gemäß Anspruch 43, umfassend:

(a) Umsetzen einer Verbindung der Formel (III):

mit einem aktivierten Derivat der 2-Furoesäure in einer Menge von mindestens 2 mol des aktivierten Derivats pro Mol der Verbindung der Formel (III), um eine Verbindung der Formel (IIA) zu erhalten:
und
(b) Entfernen des schwefelgebundenen 2-Furoylrestes von Verbindung der Formel (IIA) durch Reaktion des Produkts aus Schritt (a) mit einer organischen primären oder sekundären Aminbase, die ein wasserlösliches 2-Furoylamid bilden kann.

47. Verfahren zur Herstellung einer Verbindung der Formel (II) gemäß Anspruch 46, das ferner die folgenden Schritte umfaßt:

(c1) wenn das Produkt aus Schritt (b) in einem im wesentlichen wasserunmischbaren organischen Lösungsmittel gelöst ist, Reinigen der Verbindung der Formel (II) durch Auswaschen des Amid-Nebenprodukts aus Schritt (b) mit einer wässrigen Spülung, oder
(c2) wenn das Produkt aus Schritt (b) in einem wassermischbaren Lösungsmittel gelöst ist, Reinigen der Verbindung der Formel (II) durch Behandeln des Produkts aus Schritt (b) mit einem wässrigen Medium, um reine Verbindung der Formel (II) oder ein Salz davon auszufallen.

48. Verfahren zur Herstellung einer Verbindung der Formel (II) gemäß Anspruch 43, das folgendes umfaßt:

(a) Umsetzen einer Verbindung der Formel (III) gemäß Anspruch 46 mit einem aktivierten Derivat der 2-Furoylösäure in einer Menge von mindestens 2 mol des aktivierten Derivats pro Mol der Verbindung der Formel (III), um eine Verbindung der Formel (IIA) gemäß Anspruch 46 zu erhalten; und
(b) Entfernen des schwefelgebundenen 2-Furoylrestes von der Verbindung der Formel (IIA) durch Reaktion des Produkts aus Schritt (a) mit einem weiteren Mol der Verbindung der Formel (III), um zwei Mol der Verbindung der Formel (II) zu ergeben.

49. Verbindung der Formel (IIA):
50. Verbindung der Formel (VI):

51. Verbindung der Formel (VII):

oder ein Salz davon.

52. Verbindung der Formel (VIII):
53. Verbindung der Formel (IXA):

worin X Halogen darstellt.

54. Verbindung der Formel (IXA) gemäß Anspruch 53:

worin X Br darstellt.

55. Verbindung der Formel (XII):
oder ein Salz davon.

56. Verbindung der Formel (XV):

worin P eine Hydroxyschutzgruppe darstellt.

57. Verbindung der Formel (XVI):

58. Verbindung der Formel (XVII):
oder ein Salz davon, worin P eine Hydroxyschutzgruppe darstellt.

59. Verbindung der Formel (XX):

oder ein Salz davon oder ein Derivat, worin die 11-Carbonylgruppe maskiert ist.

60. Verbindung der Formel (XXIII):

worin L eine Abgangsgruppe, die nicht Fluor ist, darstellt.

61. Verfahren zur Herstellung einer Verbindung der Formel (I) als unsolvatisiertes Form-2-Polymorph gemäß Anspruch 4, das das Löschen der Verbindung der Formel (I) in unsolvatisierter Form in Methanol oder trockenem Dichlormethan und das Umkristallisieren der Verbindung der Formel (I) als unsolvatisiertes Form-2-Polymorph umfaßt.
62. Verfahren zur Herstellung einer Verbindung der Formel (I) als unsolvatisiertes Form-3-Polymorph gemäß Anspruch 5, das das Löschen der Verbindung der Formel (I) oder eines Solvats davon in Dichlormethan in Gegenwart von Wasser und das Umkristallisieren der Verbindung der Formel (I) als unsolvatisiertes Form-3-Polymorph umfaßt.

63. Verfahren zur Herstellung einer Verbindung der Formel (I) gemäß Anspruch 1 oder eines Solvats davon, das das Umsetzen einer Verbindung der Formel (VI):

\[
\text{(VI)}
\]

mit einer Fluorquelle umfaßt.

64. Verfahren zur Herstellung einer Verbindung der Formel (I) gemäß Anspruch 1 oder eines Solvats davon, das das Bereitstellen einer Verbindung der Formel (I), in der die 11-β-Hydroxygruppe geschützt oder maskiert ist, und das Entschützen oder Entmaskieren der Verbindung umfaßt, um die Verbindung der Formel (I) oder ein Solvat davon zu erhalten.

65. Verfahren gemäß Anspruch 64, worin die 11-β-Hydroxygruppe geschützt ist, das das Entschützen einer Verbindung der Formel (XV):

\[
\text{(XV)}
\]

umfaßt, worin \( P \) eine Hydroxyschutzgruppe darstellt.

66. Verfahren gemäß Anspruch 64, worin die 11-β-Hydroxygruppe maskiert ist, das die Reduktion einer Verbindung der Formel (XVI):
oder eines Derivats umfaßt, worin die 11-Carbonylgruppe maskiert ist.

67. Verfahren zur Herstellung einer Verbindung der Formel (I) gemäß Anspruch 1 oder eines Solvats davon, das die Reaktion einer Verbindung der Formel (XXIII):

worin L eine Abgangsgruppe darstellt, mit einer Fluorquelle umfaßt.

68. Verfahren zur Herstellung einer Verbindung der Formel (I) gemäß Anspruch 1 oder eines Solvats davon, das das Entschützen oder Entmaskieren eines Derivats einer Verbindung der Formel (I) umfaßt, in dem die 3-Carbonylgruppe geschützt oder maskiert ist.

69. Verbindung der Formel (X):
70. Verfahren zur Herstellung einer Verbindung der Formel (II) gemäß Anspruch 43, das das Behandeln einer Verbindung der Formel (X) gemäß Anspruch 69 mit einem zum Umwandeln einer Carbonsäure zu einer Thiocarbonsäure geeigneten Reagens umfaßt.

71. Verfahren zur Herstellung einer Verbindung der Formel (VI) gemäß Anspruch 50, das das Verestern einer Verbindung der Formel (IX):

![Chemical Structure]

mit einem aktivierten Derivat der 2-Furoesäure umfaßt.

72. Verfahren zur Herstellung einer Verbindung der Formel (VIII) gemäß Anspruch 52, das das Verestern einer Verbindung der Formel (XIII):

![Chemical Structure]

mit einem aktivierten Derivat der 2-Furoesäure umfaßt.

73. Verfahren zur Herstellung einer Verbindung der Formel (XII) gemäß Anspruch 55, das das Verestern einer Verbindung der Formel (XIV):
oder eines Salzes davon mit einem aktivierten Derivat der 2-Furoesäure umfaßt.

74. Verfahren zur Herstellung einer Verbindung der Formel (XVI) gemäß Anspruch 57, das das Verestern einer Verbindung der Formel (XXI):

oder eines Derivats, worin die 11-Carbonylgruppe maskiert ist, mit einem aktivierten Derivat der 2-Furoesäure umfaßt.

75. Verfahren zur Herstellung einer Verbindung der Formel (XX) gemäß Anspruch 59, das das Verestern einer Verbindung der Formel (XXII):

oder eines Derivats, worin die 11-Ketongruppe maskiert ist, mit einem aktivierten Derivat der 2-Furoesäure umfaßt.

76. Dosier inhalator, der eine pharmazeutische Aerosolformulierung gemäß Anspruch 28 umfaßt.
77. Dosierinhaltor gemäß Anspruch 76, worin die pharmazeutische Aerosolformulierung in einem unter Druck stehenden Kanister bewahrt wird, der mit einem Ventil verschlossen ist.

78. Dosierinhaltor gemäß Anspruch 76 oder 77, worin die pharmazeutische Aerosolformulierung eine Teilchengröße im Bereich von 1 bis 10 μm hat.

Revidications

1. Composé de formule (I)

2. Composé de formule (I) selon la revendication 1, sous une forme non solvatée.

3. Composé de formule (I) sous forme non solvatée selon la revendication 2, sous la form du polymorphe de Forme 1 qui a un profil de diffraction des rayons X sur poudre avec un pic à environ 18,9 degrés 2θ.

4. Composé de formule (I) sous forme non solvatée selon la revendication 2, sous la form du polymorphe de Forme 2 qui a un profil de diffraction des rayons X sur poudre avec un pic à environ 18,4 et 21,5 degrés 2θ.

5. Composé de formule (I) sous forme non solvatée selon la revendication 2, sous la form du polymorphe de Forme 3 qui a un profil de diffraction des rayons X sur poudre avec un pic à environ 18,6 et 19,2 degrés 2θ.

6. Composé de formule (I) selon la revendication 1, sous la forme d’un solide cristallin sous la forme d’un solvate essentiellement stoechiométrique avec de l’acétone.

7. Composé de formule (I) selon la revendication 1, sous la forme d’un solide cristallin sous la forme d’un solvate essentiellement stoechiométrique avec du tétrahydrofurane.

8. Composé de formule (I) selon la revendication 1, sous la forme d’un solide cristallin sous la forme d’un solvate essentiellement stoechiométrique avec l’isopropanol.

9. Composé de formule (I) selon la revendication 1, sous la forme d’un solide cristallin sous la forme d’un solvate essentiellement stoechiométrique avec de la méthyl-éthyl-cétone.

10. Composé de formule (I) selon la revendication 1, sous la forme d’un solide cristallin sous la forme d’un solvate essentiellement stoechiométrique avec du diméthylformamide.

11. Composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 destiné à une utilisation en médecine vétérinaire ou humaine.

12. Utilisation d’un composé de formule (I) ou d’un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 pour la fabrication d’un médicament destiné au traitement d’affections inflammatoires et/ou allergiques.
13. Utilisation d’un composé de formule (I) ou d’un solvate physiologiquement acceptable de celui-ci selon la revendication 12 pour le traitement d’une affection inflammatoire du nez.

14. Utilisation selon la revendication 13, où l’affection inflammatoire du nez est une rhinite.

15. Utilisation d’un composé de formule (I) ou d’un solvate physiologiquement acceptable de celui-ci selon la revendication 12 pour le traitement de la conjonctivite.

16. Utilisation selon l’une quelconque des revendications 13 à 15, où le composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci est pour une administration par inhalation.

17. Utilisation selon l’une quelconque des revendications 13 à 16, où le composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci est pour une administration par voie locale.

18. Utilisation selon l’une quelconque des revendications 13 à 16, où le composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci est pour une administration par voie topique.

19. Composition pharmaceutique comprenant un composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 en mélange avec un ou plusieurs diluants ou supports physiologiquement acceptables.

20. Composition pharmaceutique selon la revendication 19 comprenant un composé de formule (I) selon la revendication 2.

21. Composition pharmaceutique selon la revendication 19 comprenant un composé de formule (I) selon la revendication 3.

22. Composition pharmaceutique selon la revendication 19 comprenant un composé de formule (I) selon la revendication 4.

23. Composition pharmaceutique selon la revendication 19 comprenant un composé de formule (I) selon la revendication 5.

24. Composition pharmaceutique selon l’une quelconque des revendications 19 à 23 où la composition est une pulvérisation.

25. Composition pharmaceutique selon l’une quelconque des revendications 19 à 23, dans laquelle ledit composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci est présent en une quantité de 0,001 à 10 % en poids de ladite composition.

26. Formulation pharmaceutique comprenant un composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 mélangé avec un ou plusieurs diluants ou véhicules physiologiquement acceptables qui est non pressurisé et adapté pour être administrée par voie topique dans la cavité nasale.

27. Formulation pharmaceutique selon la revendication 26 qui contient de l’eau en tant que diluant ou support.

28. Formulation pharmaceutique d’aérosol comprenant un composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 et un fluorocarbure ou un chloro-fluorocarbure contenant de l’hydrogène en tant que propulseur, éventuellement en combinaison avec un surfactant et/ou un cosolvant.

29. Formulation pharmaceutique d’aérosol selon la revendication 28, qui comprend un composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 3, et un fluorocarbure ou un chloro-fluorocarbure contenant de l’hydrogène en tant que propulseur et un agent de suspension qui est soluble dans le propulseur.

30. Formulation pharmaceutique d’aérosol selon la revendication 29, dans laquelle l’agent de suspension est un acide
oligolactique ou un dérivé de celui-ci.

31. Formulation pharmaceutique d’aérosol selon l’une quelconque des revendications 28 à 30, dans laquelle le propulseur est choisi parmi le 1,1,1,2-tétrafluoroéthane, le 1,1,1,2,3,3,3-heptafluoro-n-propane et des mélanges de ceux-ci.

32. Formulation pharmaceutique d’aérosol selon la revendication 28 qui est constituée essentiellement d’un composé de formule (I) ou d’un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 3, éventuellement en combinaison avec un autre agent thérapeutiquement actif et un propulseur choisi parmi le 1,1,1,2-tétrafluoroéthane, le 1,1,1,2,3,3,3-heptafluoro-n-propane et des mélanges de ceux-ci.

33. Composition pharmaceutique selon l’une quelconque des revendications 19 à 32 qui comprend en outre un autre agent thérapeutiquement actif.

34. Composition pharmaceutique selon la revendication 33, dans laquelle ledit autre agent thérapeutiquement actif est un antihistaminique, un agent anti-inflammatoire ou un agent anti-infectieux.

35. Composition pharmaceutique selon la revendication 34, dans laquelle ledit antihistaminique est le méthapyrilène ou la loratadine, ledit agent anti-inflammatoire est un AINS et ledit agent anti-infectieux est un antibioïque ou un antiviral.

36. Composition pharmaceutique selon la revendication 33, dans laquelle ledit autre agent thérapeutiquement actif est un inhibiteur de PDE4.

37. Composition pharmaceutique selon la revendication 36, dans laquelle l’inhibiteur de PDE4 est au moins l’un choisi dans le groupe constitué de (R)\(+\)-1-(4-bromo-benzyl)-4-[(3-cyclopentoxy)-4-méthoxyphényl]-2-pyrrolidone ; 3-(cyclopentoxy-4-méthoxyphényl)-1-(4-N’-[N2-cyano-S-méthyl-isothiouréido]benzyl)-2-pyrrolidone ; acide cis-4-cyano-4-(3-cyclopentoxy-4-méthoxyphényl)cyclohexan-1-carboxylique ; cis-[4-cyano-4-(3-cyclopropylméthoxy-4-difluorométhoxyphényl)-cyclohexan-1-ol] ; [4-(3-cyclopentoxy-4-méthoxy-phényl)pyrrolidin-2-yldène]acétate de (R)\(+\)-éthyle ; [4-(3-cyclopentoxy-4-méthoxyphényl)pyrrolidin-2-yldène]acétate de (S)-(\(-\)-éthyle ; et 2-carbo-méthoxy-4-cyano-4-(3-cyclopropylméthoxy-4-difluorométhoxy-phényl)cyclohexan-1-one.

38. Composé de formule (I) ou un solvate physiologiquement acceptable de celui-ci selon l’une quelconque des revendications 1 à 5 destiné à une utilisation dans le traitement d’une affection inflammatoire et/ou allergique.

39. Procédé de préparation d’un composé de formule (I) selon la revendication 1 ou d’un solvate de celui-ci qui comprend l’alkylation du groupe 17\(\beta\)-carbothioïque d’un composé de formule (II)

ou d’un sel de celui-ci avec un composé de formule FCH\(_2\)L dans laquelle L représente un groupe libérable.

40. Procédé selon la revendication 39 dans lequel l’alkylation est réalisée en faisant réagir le composé de formule (II) ou un sel de celui-ci avec un halogénure de fluorométhyle.

41. Procédé de préparation d’un composé de formule (I) sous la forme du polymorphe de Forme 1 non solvaté selon la revendication 3 qui comprend :
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(a) la cristallisation du composé de formule (I) en présence d’un solvant non solvatant ; ou
(b) la désolvatation d’un composé de formule (I) sous une forme solvatée.

42. Procédé de préparation d’un composé de formule (I) sous la forme du polymorphe de Forme 1 non solvaté selon la revendication 3 qui comprend la dissolution du composé de formule (I) dans de la méthyl-isobutyl-cétone, de l’acétate d’éthyle ou de l’acétate de méthyle et la production du composé de formule (I) sous la Forme 1 non solvatée par addition d’un anti-solvant non solvant.

43. Composé de formule (II)

ou un sel de celui-ci.

44. Composé de formule (II) selon la revendication 43 sous la forme d’un sel cristallin solide.

45. Composé de formule (II) selon la revendication 44 sous la forme du sel de diisopropyléthylamine.

46. Procédé de préparation d’un composé de formule (II) selon la revendication 43 qui comprend :

(a) la réaction d’un composé de formule (III)

avec un dérivé activé de l’acide 2-furoïque en une quantité d’au moins 2 moles du dérivé activé par mole de composé de formule (III) pour produire un composé de formule (IIA)
et
(b) l’élimination du radical 2-furoyle lié par du soufre du composé de formule (IIA) par la réaction du produit de l’étape (a) avec une base d’amine primaire ou secondaire organique capable de former un 2-furoyl-amide hydrosoluble.

47. Procédé de préparation d’un composé de formule (II) selon la revendication 46 qui comprend en outre les étapes suivantes :

(c1) lorsque le produit de l’étape (b) est dissous dans un solvant organique pratiquement non miscible à l’eau, purifier le composé de formule (II) par élimination par lavage du sous-produit de l’amide issu de l’étape (b) avec un lavage aqueux, ou
(c2) lorsque le produit de l’étape (b) est dissous dans un solvant miscible à l’eau, purifier le composé de formule (II) par traitement du produit de l’étape (b) avec un milieu aqueux de manière à précipiter le composé de formule (II) pur ou un sel de celui-ci.

48. Procédé de préparation d’un composé de formule (II) selon la revendication 43 qui comprend :

(a) la réaction d’un composé de formule (III) selon la revendication 46 avec un dérivé activé de l’acide 2-furoïque en une quantité d’au moins 2 moles du dérivé activé par mole de composé de formule (III) pour produire un composé de formule (IIA) selon la revendication 46 ; et
(b) l’élimination du radical 2-furoyle lié par du soufre du composé de formule (IIA) par la réaction du produit de l’étape (a) avec une autre mole de composé de formule (III) pour donner deux moles de composé de formule (II).

49. Composé de formule (IIA)

50. Composé de formule (VI)
51. Composé de formule (VII)

ou un sel de celui-ci.

52. Composé de formule (VIII)

53. Composé de formule (IXA)
dans laquelle X représente un atome d'halogène.

54. Composé de formule (IXA) selon la revendication 53

dans laquelle X représente Br.

55. Composé de formule (XII)

ou un sel de celui-ci.

56. Composé de formule (XV)
dans laquelle P représente un groupe hydroxy-protecteur.

57. Composé de formule (XVI)

ou un sel de celui-ci, où P représente un groupe hydroxy-protecteur.

59. Composé de formule (XX)
ou un sel de celui-ci ou un dérivé où le groupe 11-carbonyle est masqué.

60. Composé de formule (XXIII)

\[
\text{dans laquelle } L \text{ représente un groupe libérable autre qu’un atome de fluor.}
\]

61. Procédé de préparation d’un composé de formule (I) sous la forme du polymorphe de Forme 2 non solvâté selon la revendication 4 qui comprend la dissolution du composé de formule (I) sous une forme non solvâtée dans du méthanol ou du dichlorométhane sec et la recristallisation du composé de formule (I) sous la forme du polymorphe de Forme 2 non solvâté.

62. Procédé de préparation d’un composé de formule (I) sous la forme du polymorphe de Forme 3 non solvâté selon la revendication 5 qui comprend la dissolution du composé de formule (I) ou d’un solvate de celui-ci dans du dichlorométhane en présence d’eau et la recristallisation du composé de formule (I) sous la forme du polymorphe de Forme 3 non solvâté.

63. Procédé de préparation d’un composé de formule (I) selon la revendication 1 ou d’un solvate de celui-ci qui comprend la réaction d’un composé de formule (VI)
64. Procédé de préparation d’un composé de formule (I) ou d’un solvate de celui-ci selon la revendication 1 ou d’un solvate de celui-ci qui comprend la fourniture d’un composé de formule (I) dans lequel le groupe 11-β-hydroxy est protégé ou masqué, la déprotection ou le démasquage du composé pour produire le composé de formule (I) ou un solvate de celui-ci.

65. Procédé selon la revendication 64, dans lequel le groupe 11-β-hydroxy est protégé, qui comprend la déprotection d’un composé de formule (XV)

![Image](XV)

dans laquelle P représente un groupe hydroxy-protecteur.

66. Procédé selon la revendication 64, dans lequel le groupe 11-β-hydroxy est masqué, qui comprend la réduction d’un composé de formule (XVI)

![Image](XVI)

ou d’un dérivé où le groupe 11-carbonyl est masqué.

67. Procédé de préparation d’un composé de formule (I) selon la revendication 1 ou d’un solvate de celui-ci qui comprend la réaction d’un composé de formule (XXIII)

![Image](XXIII)
dans laquelle L représente un groupe libérable avec une source de fluor.

68. Procédé de préparation d'un composé de formule (I) selon la revendication 1 ou d'un solvate de celui-ci qui comprend la déprotection ou le démasquage d'un dérivé d'un composé de formule (I) dans laquelle le groupe 3-carbonyle est protégé ou masqué.

69. Composé de formule (X)

70. Procédé de préparation d'un composé de formule (II) selon la revendication 43 qui comprend le traitement d'un composé de formule (X) selon la revendication 69 avec un réactif approprié pour la conversion d'un acide carboxylique en un acide carbothioïque.

71. Procédé de préparation d'un composé de formule (VI) selon la revendication 50 qui comprend l'estérification d'un composé de formule (IX)

avec un dérivé activé de l'acide 2-furoïque.

72. Procédé de préparation d'un composé de formule (VIII) selon la revendication 52 qui comprend l'estérification d'un composé de formule (XIII)
73. Procédé de préparation d’un composé de formule (XII) selon la revendication 55 qui comprend l’estérisation d’un composé de formule (XIV)

ou d’un sel de celui-ci, avec un dérivé activé de l’acide 2-furoïque.

74. Procédé de préparation d’un composé de formule (XVI) selon la revendication 57 qui comprend l’estérisation d’un composé de formule (XXI)

ou d’un dérivé où le groupe 11-carbonyle est masqué, avec un dérivé activé de l’acide 2-furoïque.

75. Procédé de préparation d’un composé de formule (XX) selon la revendication 59 qui comprend l’estérisation d’un composé de formule (XXII)

ou d’un dérivé où le groupe 11-cétone est masqué, avec un dérivé activé de l’acide 2-furoïque.

76. Inhalateur doseur comprenant une formulation pharmaceutique d’aérosol selon la revendication 28.
77. Inhalateur doseur selon la revendication 76, dans lequel ladite formulation pharmaceutique d’aérosol est gardée dans un récipient métallique scellé pressurisé fermé par une valve.

78. Inhalateur doseur selon la revendication 76 ou la revendication 77, dans lequel ladite formulation pharmaceutique a une taille de particule située dans la plage de 1 à 10 μm.
REFERENCES CITED IN THE DESCRIPTION

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